216

Attorney Docket: PATH03-16

Express Mail Label:

EL442001728 US

Date: December 1,2003

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Title Of The Invention

NUCLEIC ACID AND AMINO ACID SEQUENCES RELATING TO STAPHYLOCOCCUS EPIDERMIDIS FOR DIAGNOSTICS AND THERAPEUTICS

Related Applications

This application is a Continuation-in-Part which claims priority to U.S. Application No. 09/134,001, filed August 13, 1998, which claims the benefit of both U.S. Provisional Application No. 60/064,964, filed November 8, 1997 and U.S. Provisional Application No. 60/055,779, filed August 14, 1997, the contents of all of which are incorporated in their entirety.

Background Of The Invention

that are Gram-positive, nonmotile, nonpigmented and coagulase-negative cocci, which are mainly found on the skin and mucous membrane of warm-blooded animals. Their large numbers and ubiquitous distribution result in frequent contamination of specimens collected from or through the skin, making these organisms amongst the most frequently isolated in the clinical laboratory. In the past, *S. epidermidis* was rarely the cause of significant infections, but with the increasing use of implanted catheters and prosthetic devices, it has emerged as an important agent of hospital-acquired infections and has been recognized as a true pathogen (Lowy and Hammer, 1983, Ann Intern Med, <u>99</u>: 834-9; Blum and Rodvold, 1987, Clin Pharm, <u>6</u>: 464-75; Hamory, Parisi et al., 1987, Am J Infect Control, <u>15</u>: 59-74). *S. epidermidis* is a major cause of infection of indwelling foreign devices such as, orthopedic devices, intravenous catheters, prosthetic heart valves, central nervous system shunts, and peritoneal dialysis catheters (Blum and Rodvold, 1987, Clin Pharm, <u>6</u>: 464-75; Archer, 1988,

J Antimicrob Chemother, <u>21 Suppl C</u>: 133-8)(Lowy and Hammer, 1983, Ann Intern Med, <u>99</u>: 834-9; Hamory, Parisi et al., *Staphylococcus* 1987, Am J Infect Control, <u>15</u>: 59-74). In addition *S. epidermidis* is a common cause of postoperative wound infections, bacteremia of immunosuppressed patients, intensive-care unit patients and premature newborns (MacLowry, 1983, Am J Med, <u>75</u>: 2-6)(Eykyn, 1988, Lancet, <u>1</u>: 100-4). According to a national survey (Centers for Disease Control, 1981:7) *S. epidermidis* caused 8.9% of primary nosocomial bacteremias.

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Treatment of S. epidermidis infections remains difficult because of the occult nature, association with foreign bodies, and frequent resistance to antimicrobial agents. Ordinarily, S. epidermidis is an organism with low virulence, however breaks in host defense caused by surgery, catheter placement, prosthesis insertion or immuno-suppression is prerequisite for infection. The presence of foreign bodies itself facilitates infection by protecting the organism from elimination by host defenses or antimicrobial therapy (Lowy and Hammer, 1983, Ann Intern Med, 99: 834-9). Furthermore, S. epidermidis due to its ability to produce extracellular polysaccharide material or slime, may be uniquely adapted to adhere to smooth surfaces such as plastics or metal. Slime producing strains of S. epidermidis appear to be more pathogenic than non-slime producing strains (Christensen, Simpson et al., 1983, Infect Immun, 40: 407-10; Peters and Pulverer, 1984, J Antimicrob Chemother, 14 Suppl D: 67-71; Gallimore, Gagnon et al., 1991, J Infect Dis, 164: 1220-3). This property and many factors are involved in the pathogenesis of device associated infections. Despite the increased recognition as a pathogen, S. epidermidis infections are difficult to diagnose. Differentiating clinically important from clinically unimportant bacterial isolates of S. epidermidis is difficult because of the high rate of contamination.

Although laboratory isolates of *S. epidermidis* have generally been susceptible to semisynthetic penicillins (methicillin, nafcillin, oxacillin), cephalosporins, amino-glycosides, vancomycin and rafampin, recent clinical isolates have had an increased resistance. Recent reports (Karchmer, 1985, Am J Med, <u>78</u>: 116-27; Karchmer, 1991, J Hosp Infect, <u>18 Suppl</u> <u>A</u>: 355-66) show that 83% of *S. epidermidis* isolates from patients with prosthetic valve endocarditis are methicillin resistant and 32% are gentamicin resistant as well. Multi-drug resistant staphylococci have emerged in the midst of high level use of penicillin and

aminoglycosides (Centers for Disease Control and Prevention, 1993 *MMWR* 42:597; and S. Handwerger et al., 1993, *Clin Infect Dis* 16:750).

The use of antibiotics for therapeutics and prophylactic purposes, promotes the selection of resistant organisms and the spread of antibiotic resistance genes among bacteria. Previous studies have shown that virtually all staphylococci carry some antibiotic resistance genes on naturally occurring extrachromosomal mobile genetic elements, such as the plasmids. Survey and analysis of plasmids in clinical isolates of S. epidermidis have shown that more that 80% of isolates carry plasmids and in several cases more than one plasmid (Archer et al., 1982, Infect Immun, 35:627-632; Kloos et al., 1981, Can J Microbiol, 27:271-278; Moller, 1988, J Hosp Infect 12:19-27). Though the most important forms of resistance has been the inactivation of antibiotics, particularly penicillins and cephalosporins, recent clinical isolates have resistance to one or more of the following antibiotics, methicillin, tetracycline, erythromycin, gentamycin, kanamycin and chloramphenicol. In fact due to the wide spread occurrence of plasmids and their involvement in antibiotic resistance, plasmid profiling has been used as an epidemiological reagent to study nosocomial infections. This invention relates to isolated nucleic acids and polypeptides derived from S. epidermidis plasmids that are useful as molecular targets for diagnosis, prophylaxis and treatment of pathological conditions, as well as materials and methods for the diagnosis, prevention, and amelioration of pathological conditions resulting from bacterial infection.

These concerns point to the need for diagnostic tools and therapeutics aimed at proper identification of strain and eradication of virulence. The design of vaccines that will limit the spread of infection and halt transfer of resistance factors is very desirable.

Summary Of The Invention

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The present invention fulfills the need for diagnostic tools and therapeutics by providing bacterial-specific compositions and methods for detecting *Staphylococcus* species including *S. epidermidis*, as well as compositions and methods useful for treating and preventing *Staphylococcus* infection, in particular, *S. epidermidis* infection, in vertebrates including mammals.

The present invention encompasses isolated nucleic acids and polypeptides derived from *S. epidermidis* that are useful as reagents for diagnosis of bacterial disease, components of effective antibacterial vaccines, and/or as targets for antibacterial drugs including anti-*S. epidermidis* drugs. They can also be used to detect the presence of *S. epidermidis* and other *Staphylococcus* species in a sample; and in screening compounds for the ability to interfere with the *S. epidermidis* life cycle or to inhibit *S. epidermidis* infection. They also have use as biocontrol agents for plants.

In one aspect, the invention features compositions of nucleic acids corresponding to entire coding sequences of *S. epidermidis* proteins (SEQ ID NO: 1 - SEQ ID NO: 3702), including surface or secreted proteins or parts thereof, nucleic acids capable of binding mRNA from *S. epidermidis* proteins to block protein translation, and methods for producing *S. epidermidis* proteins or parts thereof using peptide synthesis and recombinant DNA techniques. This invention also features antibodies and nucleic acids useful as probes to detect *S. epidermidis* infection. In addition, vaccine compositions and methods for the protection or treatment of infection by *S. epidermidis* are within the scope of this invention.

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In another aspect, the invention relates to the nucliec acids corresponding to 2 naturally occurring plasmids of *S. epidermidis* (SEQ ID NO: 3703 - SEQ ID NO: 3772) and the corresponding amino acid sequences (SEQ ID NO: 7475 - SEQ ID NO: 7544).

The nucleotide sequences provided in SEQ ID NO: 1 - SEQ ID NO: 3772, a fragment thereof, or a nucleotide sequence at least about 99.5% identical to a sequence contained within SEQ ID NO: 1 - SEQ ID NO: 3772 may be "provided" in a variety of medias to facilitate use thereof. As used herein, "provided" refers to a manufacture, other than an isolated nucleic acid molecule, which contains a nucleotide sequence of the present invention, i.e., the nucleotide sequence provided in SEQ ID NO: 1 - SEQ ID NO: 3772, a fragment thereof, or a nucleotide sequence at least about 99.5% identical to a sequence contained within SEQ ID NO: 1 - SEQ ID NO: 3772. Uses for and methods for providing nucleotide sequences in a variety of media is well known in the art (see e.g., EPO Publication No. EP 0 756 006).

In one application of this embodiment, a nucleotide sequence of the present invention can be recorded on computer readable media. As used herein, "computer readable media" refers to any media which can be read and accessed directly by a computer. Such media include, but are not limited to: magnetic storage media, such as floppy discs, hard disc storage media, and magnetic tape; optical storage media such as CD-ROM; electrical storage media such as RAM and ROM; and hybrids of these categories such as magnetic/optical storage media. A person skilled in the art can readily appreciate how any of the presently known computer readable media can be used to create a manufacture comprising computer readable media having recorded thereon a nucleotide sequence of the present invention.

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As used herein, "recorded" refers to a process for storing information on computer readable media. A person skilled in the art can readily adopt any of the presently known methods for recording information on computer readable media to generate manufactures comprising the nucleotide sequence information of the present invention.

A variety of data storage structures are available to a person skilled in the art for creating a computer readable media having recorded thereon a nucleotide sequence of the present invention. The choice of the data storage structure will generally be based on the means chosen to access the stored information. In addition, a variety of data processor programs and formats can be used to store the nucleotide sequence information of the present invention on computer readable media. The sequence information can be represented in a word processing text file, formatted in commercially-available software such as WordPerfect and Microsoft Word, or represented in the form of an ASCII file, stored in a database application, such as DB2, Sybase, Oracle, or the like. A person skilled in the art can readily adapt any number of data processor structuring formats (e.g. text file or database) in order to obtain computer readable media having recorded thereon the nucleotide sequence information of the present invention.

By providing the nucleotide sequence of SEQ ID NO: 1 - SEQ ID NO: 3772, a fragment thereof, or a nucleotide sequence at least about 99.5% identical to SEQ ID NO: 1 -

SEQ ID NO: 3772 in computer readable form, a person skilled in the art can routinely access the coding sequence information for a variety of purposes. Computer software is publicly available which allows a person skilled in the art to access sequence information provided in a computer readable media. Examples of such computer software include programs of the "Staden Package", "DNA Star", "MacVector", GCG "Wisconsin Package" (Genetics Computer Group, Madison, WI) and "NCBI Toolbox" (National Center For Biotechnology Information). Suitable programs are described, for example, in Martin J. Bishop, ed., *Guide to Human Genome Computing*, 2d Edition, Academic Press, San Diego, CA. (1998); and Leonard F. Peruski, Jr., and Anne Harwood Peruski, *The Internet and the New Biology: Tools for Genomic and Molecular Research*, American Society for Microbiology, Washington, D.C. (1997).

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Computer algorithms enable the identification of S. epidermidis open reading frames (ORFs) within SEQ ID NO: 1 - SEQ ID NO: 3772 which contain homology to ORFs or proteins from other organisms. Examples of such similarity-search algorithms include the 15 BLAST [Altschul et al., J. Mol. Biol. 215:403-410 (1990)] and Smith-Waterman [Smith and Waterman (1981) Advances in Applied Mathematics, 2:482-489] search algorithms. Suitable search algorithms are described, for example, in Martin J. Bishop, ed., Guide to Human Genome Computing, 2d Edition, Academic Press, San Diego, CA. (1998); and Leonard F. Peruski, Jr., and Anne Harwood Peruski, The Internet and the New Biology: Tools for Genomic and Molecular Research, American Society for Microbiology, Washington, D.C. 20 (1997). Such algorithms are utilized on computer systems as exemplified below. The ORFs so identified represent protein encoding fragments within the S. epidermidis genome and S. epidermidis plasmidsand are useful in producing commercially important proteins such as enzymes used in fermentation reactions and in the production of commercially useful metabolites. 25

The present invention further provides systems, particularly computer-based systems, which contain the sequence information described herein. Such systems are designed to

identify commercially important fragments of the *S. epidermidis* genome and plasmids. As used herein, "a computer-based system" refers to the hardware means, software means, and data storage means used to analyze the nucleotide sequence information of the present invention. The minimum hardware means of the computer-based systems of the present invention comprises a central processing unit (CPU), input means, output means, and data storage means. A person skilled in the art can readily appreciate that any one of the currently available computer-based systems is suitable for use in the present invention. The computer-based systems of the present invention comprise a data storage means having stored therein a nucleotide sequence of the present invention and the necessary hardware means and software means for supporting and implementing a search means. As used herein, "data storage means" refers to memory which can store nucleotide sequence information of the present invention, or a memory access means which can access manufactures having recorded thereon the nucleotide sequence information of the present invention.

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As used herein, "search means" refers to one or more programs which are 15 implemented on the computer-based system to compare a target sequence or target structural motif with the sequence information stored within the data storage means. Search means are used to identify fragments or regions of the S. epidermidis genome and plasmids which are similar to, or "match", a particular target sequence or target motif. A variety of known algorithms are known in the art and have been disclosed publicly, and a variety of 20 commercially available software for conducting homology-based similarity searches are available and can be used in the computer-based systems of the present invention. Examples of such software includes, but is not limited to, FASTA (GCG Wisconsin Package), Bic SW (Compugen Bioccelerator), BLASTN2, BLASTP2, BLASTX2 (NCBI) and Motifs (GCG). Suitable software programs are described, for example, in Martin J. Bishop, ed., Guide to Human Genome Computing, 2d Edition, Academic Press, San Diego, CA. (1998); and 25 Leonard F. Peruski, Jr., and Anne Harwood Peruski, The Internet and the New Biology: Tools for Genomic and Molecular Research, American Society for Microbiology,

Washington, D.C. (1997). A person skilled in the art can readily recognize that any one of the available algorithms or implementing software packages for conducting homology searches can be adapted for use in the present computer-based systems.

As used herein, a "target sequence" can be any DNA or amino acid sequence of six or more nucleotides or two or more amino acids. A person skilled in the art can readily recognize that the longer a target sequence is, the less likely a target sequence will be present as a random occurrence in the database. The most preferred sequence length of a target sequence is from about 10 to 100 amino acids or from about 30 to 300 nucleotide residues. However, it is well recognized that many genes are longer than 500 amino acids, or 1.5 kb in length, and that commercially important fragments of the *S. epidermidis* genome and plasmids from *S. epidermidis*, such as sequence fragments involved in gene expression and protein processing, will often be shorter than 30 nucleotides.

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As used herein, "a target structural motif," or "target motif," refers to any rationally selected sequence or combination of sequences in which the sequence(s) are chosen based on a specific functional domain or three-dimensional configuration which is formed upon the folding of the target polypeptide. There are a variety of target motifs known in the art. Protein target motifs include, but are not limited to, enzymatic active sites, membrane-spanning regions, and signal sequences. Nucleic acid target motifs include, but are not limited to, promoter sequences, hairpin structures and inducible expression elements (protein binding sequences).

A variety of structural formats for the input and output means can be used to input and output the information in the computer-based systems of the present invention. A preferred format for an output means ranks fragments of the *S. epidermidis* genome and plasmids possessing varying degrees of homology to the target sequence or target motif. Such presentation provides a person skilled in the art with a ranking of sequences which contain various amounts of the target sequence or target motif and identifies the degree of homology contained in the identified fragment.

A variety of comparing means can be used to compare a target sequence or target motif with the data storage means to identify sequence fragments of the *S. epidermidis* genome and plasmids. In the present examples, implementing software which implement the BLASTP2 and bic_SW algorithms (Altschul et al., J Mol. Biol. 215:403-410 (1990);

Compugen Biocellerator) was used to identify open reading frames within the *S. epidermidis* genome and plasmids. A person skilled in the art can readily recognize that any one of the publicly available homology search programs can be used as the search means for the computer-based systems of the present invention. Suitable programs are described, for example, in Martin J. Bishop, ed., *Guide to Human Genome Computing*, 2d Edition,
Academic Press, San Diego, CA. (1998); and Leonard F. Peruski, Jr., and Anne Harwood Peruski, *The Internet and the New Biology: Tools for Genomic and Molecular Research*, American Society for Microbiology, Washington, D.C. (1997).

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The invention features *S. epidermidis* polypeptides, preferably a substantially pure preparation of an *S. epidermidis* polypeptide, or a recombinant *S. epidermidis* polypeptide. In preferred embodiments: the polypeptide has biological activity; the polypeptide has an amino acid sequence at least about 60%, 70%, 80%, 90%, 95%, 98%, or 99% identical to an amino acid sequence of the invention contained in the Sequence Listing, preferably it has about 65% sequence identity with an amino acid sequence of the invention contained in the Sequence Listing, and most preferably it has about 92% to about 99% sequence identity with an amino acid sequence of the invention contained in the Sequence Listing; the polypeptide has an amino acid sequence essentially the same as an amino acid sequence of the invention contained in the Sequence Listing; the polypeptide is at least about 5, 10, 20, 50, 100, or 150 amino acid residues in length; the polypeptide includes at least about 5, preferably at least about 10, more preferably at least about 20, still more preferably at least about 50, 100, or 150 contiguous amino acid residues of the invention contained in the Sequence Listing. In yet another preferred embodiment, the amino acid sequence which differs in sequence identity by

about 7% to about 8% from the *S. epidermidis* amino acid sequences of the invention contained in the Sequence Listing is also encompassed by the invention.

In preferred embodiments: the *S. epidermidis* polypeptide is encoded by a nucleic acid of the invention contained in the Sequence Listing, or by a nucleic acid having at least about 60%, 70%, 80%, 90%, 95%, 98%, or 99% homology with a nucleic acid of the invention contained in the Sequence Listing.

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In a preferred embodiment, the subject *S. epidermidis* polypeptide differs in amino acid sequence at about 1, 2, 3, 5, 10 or more residues from a sequence of the invention contained in the Sequence Listing. The differences, however, are such that the *S. epidermidis* polypeptide exhibits an *S. epidermidis* biological activity, e.g., the *S. epidermidis* polypeptide retains a biological activity of a naturally occurring *S. epidermidis* enzyme.

In preferred embodiments, the polypeptide includes all or a fragment of an amino acid sequence of the invention contained in the Sequence Listing; fused, in reading frame, to additional amino acid residues, preferably to residues encoded by genomic DNA 5' or 3' to the genomic DNA which encodes a sequence of the invention contained in the Sequence Listing.

In yet other preferred embodiments, the *S. epidermidis* polypeptide is a recombinant fusion protein having a first *S. epidermidis* polypeptide portion and a second polypeptide portion, e.g., a second polypeptide portion having an amino acid sequence unrelated to *S. epidermidis*. The second polypeptide portion can be, e.g., any of glutathione-S-transferase, a DNA binding domain, or a polymerase activating domain. In preferred embodiment the fusion protein can be used in a two-hybrid assay.

Polypeptides of the invention include those which arise as a result of alternative transcription events, alternative RNA splicing events, and alternative translational and postranslational events.

In a preferred embodiment, the encoded *S. epidermidis* polypeptide differs (e.g., by amino acid substitution, addition or deletion of at least one amino acid residue) in amino acid sequence at about 1, 2, 3, 5, 10 or more residues, from a sequence of the invention contained

in the Sequence Listing. The differences, however, are such that: the *S. epidermidis* encoded polypeptide exhibits an *S. epidermidis* biological activity, e.g., the encoded *S. epidermidis* enzyme retains a biological activity of a naturally occurring *S. epidermidis*.

In preferred embodiments, the encoded polypeptide includes all or a fragment of an amino acid sequence of the invention contained in the Sequence Listing; fused, in reading frame, to additional amino acid residues, preferably to residues encoded by genomic DNA 5' or 3' to the genomic DNA which encodes a sequence of the invention contained in the Sequence Listing.

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The *S. epidermidis* strain, 18972, from which genomic sequences have been sequenced, has been deposited on July 10, 1997, in the American Type Culture Collection and assigned the ATCC designation # 55998.

Included in the invention are: allelic variations; natural mutants; induced mutants; proteins encoded by DNA that hybridize under high or low stringency conditions to a nucleic acid which encodes a polypeptide of the invention contained in the Sequence Listing (for definitions of high and low stringency see Current Protocols in Molecular Biology, John Wiley & Sons, New York, 1989, 6.3.1 - 6.3.6, hereby incorporated by reference); and, polypeptides specifically bound by antisera to *S. epidermidis* polypeptides, especially by antisera to an active site or binding domain of *S. epidermidis* polypeptide. The invention also includes fragments, preferably biologically active fragments. These and other polypeptides are also referred to herein as *S. epidermidis* polypeptide analogs or variants.

The invention further provides nucleic acids, e.g., RNA or DNA, encoding a polypeptide of the invention. This includes double stranded nucleic acids as well as coding and antisense single strands.

In preferred embodiments, the subject *S. epidermidis* nucleic acid will include a transcriptional regulatory sequence, e.g., at least one of a transcriptional promoter or transcriptional enhancer sequence, operably linked to the *S. epidermidis* gene sequence, e.g., to render the *S. epidermidis* gene sequence suitable for expression in a recombinant host cell.

In yet a further preferred embodiment, the nucleic acid which encodes an *S. epidermidis* polypeptide of the invention, hybridizes under stringent conditions to a nucleic acid probe corresponding to at least about 8 consecutive nucleotides of the invention contained in the Sequence Listing; more preferably to at least about 12 consecutive nucleotides of the invention contained in the Sequence Listing; still more preferably to at least about 20 consecutive nucleotides of the invention contained in the Sequence Listing; most preferably to at least about 40 consecutive nucleotides of the invention contained in the Sequence Listing.

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In another aspect, the invention provides a substantially pure nucleic acid having a nucleotide sequence which encodes an *S. epidermidis* polypeptide. In preferred embodiments: the encoded polypeptide has biological activity; the encoded polypeptide has an amino acid sequence at least about 60%, 70%, 80%, 90%, 95%, 98% or 99% homologous to an amino acid sequence of the invention contained in the Sequence Listing; the encoded polypeptide has an amino acid sequence essentially the same as an amino acid sequence of the invention contained in the Sequence Listing; the encoded polypeptide is at least about 5, 10, 20, 50, 100, or 150 amino acids in length; the encoded polypeptide comprises at least about 5, preferably at least about 10, more preferably at least about 20, still more preferably at least about 50, 100, or 150 contiguous amino acids of the invention contained in the Sequence Listing.

In another aspect, the invention encompasses: a vector including a nucleic acid which encodes an *S. epidermidis* polypeptide or an *S. epidermidis* polypeptide variant as described herein; a host cell transfected with the vector; and a method of producing a recombinant *S. epidermidis* polypeptide or *S. epidermidis* polypeptide variant; including culturing the cell, e.g., in a cell culture medium, and isolating *an S. epidermidis* or *S. epidermidis* polypeptide variant, e.g., from the cell or from the cell culture medium.

One embodiment of the invention is directed to substantially isolated nucleic acids.

Nucleic acids of the invention include sequences comprising at least about 8 nucleotides in

length, more preferably at least about 12 nucleotides in length, even more preferably at least about 15-20 nucleotides in length, that correspond to a subsequence of any one of SEQ ID NO: 1 - SEQ ID NO: 3772 or complements thereof. Alternatively, the nucleic acids comprise sequences contained within any ORF (open reading frame), including a complete protein-coding sequence, of which any of SEQ ID NO: 1 - SEQ ID NO: 3772 forms a part. The invention encompasses sequence-conservative variants and function-conservative variants of these sequences. The nucleic acids may be DNA, RNA, DNA/RNA duplexes, protein-nucleic acid (PNA), or derivatives thereof.

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In another aspect, the invention features a purified recombinant nucleic acid having at least about 50%, 60%, 70%, 80%, 90%, 95%, 98%, or 99% homology with a sequence of the invention contained in the Sequence Listing

The invention also encompasses recombinant DNA (including DNA cloning and expression vectors) comprising these *S. epidermidis* -derived sequences; host cells comprising such DNA, including fungal, bacterial, yeast, plant, insect, and mammalian host cells; and methods for producing expression products comprising RNA and polypeptides encoded by the *S. epidermidis* sequences. These methods are carried out by incubating a host cell comprising an *S. epidermidis* -derived nucleic acid sequence under conditions in which the sequence is expressed. The host cell may be native or recombinant. The polypeptides can be obtained by (a) harvesting the incubated cells to produce a cell fraction and a medium fraction; and (b) recovering the *S. epidermidis* polypeptide from the cell fraction, the medium fraction, or both. The polypeptides can also be made by *in vitro* translation.

In another aspect, the invention features nucleic acids capable of binding mRNA of *S. epidermidis*. Such nucleic acid is capable of acting as antisense nucleic acid to control the translation of mRNA of *S. epidermidis*. A further aspect features a nucleic acid which is capable of binding specifically to an *S. epidermidis* nucleic acid. These nucleic acids are also referred to herein as complements and have utility as probes and as capture reagents.

In another aspect, the invention features an expression system comprising an open reading frame corresponding to *S. epidermidis* nucleic acid. The nucleic acid further comprises a control sequence compatible with an intended host. The expression system is useful for making polypeptides corresponding to *S. epidermidis* nucleic acid.

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In another aspect, the invention encompasses: a vector including a nucleic acid which encodes an *S. epidermidis* polypeptide or an *S. epidermidis* polypeptide variant as described herein; a host cell transfected with the vector; and a method of producing a recombinant *S. epidermidis* polypeptide or *S. epidermidis* polypeptide variant; including culturing the cell, e.g., in a cell culture medium, and isolating the *S. epidermidis* or *S. epidermidis* polypeptide variant, e.g., from the cell or from the cell culture medium.

In yet another embodiment of the invention encompasses reagents for detecting bacterial infection, including *S. epidermidis* infection, which comprise at least one *S. epidermidis* -derived nucleic acid defined by any one of SEQ ID NO: 1 - SEQ ID NO: 3772, or sequence-conservative or function-conservative variants thereof. Alternatively, the diagnostic reagents comprise nucleotide sequences that are contained within any open reading frames (ORFs), including preferably complete protein-coding sequences, contained within any of SEQ ID NO: 1 - SEQ ID NO: 3772, or polypeptide sequences contained within any of SEQ ID NO: 3773 - SEQ ID NO: 7544, or polypeptides of which any of the above sequences forms a part, or antibodies directed against any of the above peptide sequences or function-conservative variants and/or fragments thereof.

The invention further provides antibodies, preferably monoclonal antibodies, which specifically bind to the polypeptides of the invention. Methods are also provided for producing antibodies in a host animal. The methods of the invention comprise immunizing an animal with at least one *S. epidermidis* -derived immunogenic component, wherein the immunogenic component comprises one or more of the polypeptides encoded by any one of SEQ ID NO: 1 - SEQ ID NO: 3772 or sequence-conservative or function-conservative variants thereof; or polypeptides that are contained within any ORFs, including complete

protein-coding sequences, of which any of SEQ ID NO: 1 - SEQ ID NO: 3772 forms a part; or polypeptide sequences contained within any of SEQ ID NO: 3773 - SEQ ID NO: 7544; or polypeptides of which any of SEQ ID NO: 3773 - SEQ ID NO: 7544 forms a part. Host animals include any warm blooded animal, including without limitation mammals and birds. Such antibodies have utility as reagents for immunoassays to evaluate the abundance and distribution of *S. epidermidis* -specific antigens.

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In yet another aspect, the invention provides diagnostic methods for detecting *S. epidermidis* antigenic components or anti-*S. epidermidis* antibodies in a sample. *S. epidermidis* antigenic components may be detected by known processes, including but not limited to detection by a process comprising: (i) contacting a sample suspected to contain a bacterial antigenic component with a bacterial-specific antibody, under conditions in which a stable antigen-antibody complex can form between the antibody and bacterial antigenic components in the sample; and (ii) detecting any antigen-antibody complex formed in step (i), wherein detection of an antigen-antibody complex indicates the presence of at least one bacterial antigenic component in the sample. In different embodiments of this method, the antibodies used are directed against a sequence encoded by any of SEQ ID NO: 1 - SEQ ID NO: 3772 or sequence-conservative or function-conservative variants thereof, or against a polypeptide sequence contained in any of SEQ ID NO: 3773 - SEQ ID NO: 7544 or function-conservative variants thereof.

In yet another aspect, the invention provides a method for detecting antibacterial-specific antibodies in a sample, which comprises: (i) contacting a sample suspected to contain antibacterial-specific antibodies with an *S. epidermidis* antigenic component, under conditions in which a stable antigen-antibody complex can form between the *S. epidermidis* antigenic component and antibacterial antibodies in the sample; and (ii) detecting any antigenantibody complex formed in step (i), wherein detection of an antigen-antibody complex indicates the presence of antibacterial antibodies in the sample. In different embodiments of this method, the antigenic component is encoded by a sequence contained in any of SEQ ID

NO: 1 - SEQ ID NO: 3772 or sequence-conservative and function-conservative variants thereof, or is a polypeptide sequence contained in any of SEQ ID NO: 3773 - SEQ ID NO: 7544 or function-conservative variants thereof.

In another aspect, the invention features a method of generating vaccines for immunizing an individual against *S. epidermidis*. The method includes: immunizing a subject with an *S. epidermidis* polypeptide, e.g., a surface or secreted polypeptide, or a combination of such peptides or active portion(s) thereof, and a pharmaceutically acceptable carrier. Such vaccines have therapeutic and prophylactic utilities.

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In another aspect, the invention features a method of evaluating a compound, e.g., a polypeptide, e.g., a fragment of a host cell polypeptide, for the ability to bind an *S*. *epidermidis* polypeptide. The method includes contacting the compound to be evaluated with an *S. epidermidis* polypeptide and determining if the compound binds or otherwise interacts with the *S. epidermidis* polypeptide. Compounds which bind or otherwise interact with *S. epidermidis* polypeptides are candidates as modulators, including activators and inhibitors, of the bacterial life cycle. These assays can be performed *in vitro* or *in vivo*.

In another aspect, the invention features a method of evaluating a compound, e.g., a polypeptide, e.g., a fragment of a host cell polypeptide, for the ability to bind an *S. epidermidis* nucleic acid, e.g., DNA or RNA. The method includes contacting the compound to be evaluated with an *S. epidermidis* nucleic acid and determining if the compound binds or otherwise interacts with the *S. epidermidis* nucleic acid. Compounds which bind *S. epidermidis* are candidates as modultors, including activators and inhibitors, of the bacterial life cycle. These assays can be performed *in vitro* or *in vivo*.

A particularly preferred embodiment of the invention is directed to a method of screening test compounds for anti-bacterial activity, which method comprises: selecting as a target a bacterial specific sequence, which sequence is essential to the viability of a bacterial species; contacting a test compound with said target sequence; and selecting those test compounds which bind to said target sequence as potential anti-bacterial candidates. In one

embodiment, the target sequence selected is specific to a single species, or even a single strain, such as, for example, the strain *S. epidermidis* 18972. In a second embodiment, the target sequence is common to at least two species of bacteria. In a third embodiment, the target sequence is common to a family of bacteria. The target sequence may be a nucleic acid sequence or a polypeptide sequence. Methods employing sequences common to more than one species of microorganism may be used to screen candidates for broad spectrum antibacterial activity.

The invention also provides methods for preventing or treating disease caused by certain bacteria, including *S. epidermidis*, which are carried out by administering to an animal in need of such treatment, in particular a warm-blooded vertebrate, including but not limited to birds and mammals, a compound that specifically inhibits or interferes with the function of a bacterial polypeptide or nucleic acid. In a particularly preferred embodiment, the mammal to be treated is human.

15 DETAILED DESCRIPTION OF THE INVENTION

The sequences of the present invention include the specific nucleic acid and amino acid sequences set forth in the Sequence Listing that forms a part of the present specification, and which are designated SEQ ID NO: 1 - SEQ ID NO: 7544. Use of the terms "SEQ ID NO: 1 - SEQ ID NO: 3772 ", " SEQ ID NO: 3773 - SEQ ID NO: 7544, "the sequences depicted in Table 2", etc., is intended, for convenience, to refer to each individual SEQ ID NO *individually*, and is not intended to refer to the genus of these sequences unless such reference would be indicated. In other words, it is a shorthand for listing all of these sequences individually. The invention encompasses each sequence individually, as well as any combination thereof.

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Definitions

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"Nucleic acid" or "polynucleotide" as used herein refers to purine- and pyrimidine-containing polymers of any length, either polyribonucleotides or polydeoxyribonucleotides or mixed polyribo-polydeoxyribo nucleotides. This includes single- and double-stranded molecules, i.e., DNA-DNA, DNA-RNA and RNA-RNA hybrids, as well as "protein nucleic acids" (PNA) formed by conjugating bases to an amino acid backbone. This also includes nucleic acids containing modified bases.

A nucleic acid or polypeptide sequence that is "derived from" a designated sequence refers to a sequence that corresponds to a region of the designated sequence. For nucleic acid 10 sequences, this encompasses sequences that are homologous or complementary to the sequence, as well as "sequence-conservative variants" and "function-conservative variants." For polypeptide sequences, this encompasses "function-conservative variants." Sequenceconservative variants are those in which a change of one or more nucleotides in a given codon position results in no alteration in the amino acid encoded at that position. Functionconservative variants are those in which a given amino acid residue in a polypeptide has been 15 changed without altering the overall conformation and function of the native polypeptide, including, but not limited to, replacement of an amino acid with one having similar physicochemical properties (such as, for example, acidic, basic, hydrophobic, and the like). "Function-conservative" variants also include any polypeptides that have the ability to elicit 20 antibodies specific to a designated polypeptide.

An "S. epidermidis -derived" nucleic acid or polypeptide sequence may or may not be present in other bacterial species, and may or may not be present in all S. epidermidis strains. This term is intended to refer to the source from which the sequence was originally isolated. Thus, an S. epidermidis -derived polypeptide, as used herein, may be used, e.g., as a target to screen for a broad spectrum antibacterial agent, to search for homologous proteins in other species of bacteria or in eukaryotic organisms such as bacteria humans, etc.

A purified or isolated polypeptide or a substantially pure preparation of a polypeptide are used interchangeably herein and, as used herein, mean a polypeptide that has been separated from other proteins, lipids, and nucleic acids with which it naturally occurs. Preferably, the polypeptide is also separated from substances, e.g., antibodies or gel matrix, e.g., polyacrylamide, which are used to purify it. Preferably, the polypeptide constitutes at least about 10, 20, 50 70, 80 or 95% dry weight of the purified preparation. Preferably, the preparation contains sufficient polypeptide to allow protein sequencing; at least about 1, 10, or preferably 100 mg of polypeptide.

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A purified preparation of cells refers to, in the case of plant or animal cells, an *in vitro* preparation of cells and not an entire intact plant or animal. In the case of cultured cells or microbial cells, it consists of a preparation of at least about 10%, more preferably at least about 50%, of the subject cells.

A purified or isolated or a substantially pure nucleic acid, e.g., a substantially pure DNA, (are terms used interchangeably herein) is a nucleic acid which is one or both of the following: not immediately contiguous with both of the coding sequences with which it is immediately contiguous (i.e., one at the 5' end and one at the 3' end) in the naturally-occurring genome and plasmids of the organism from which the nucleic acid is derived; or which is substantially free of a nucleic acid with which it occurs in the organism from which the nucleic acid is derived. The term includes, for example, a recombinant DNA which is incorporated into a vector, e.g., into an autonomously replicating plasmid or virus, or into the genomic DNA of a prokaryote or eukaryote, or which exists as a separate molecule (e.g., a cDNA or a genomic DNA fragment produced by PCR or restriction endonuclease treatment) independent of other DNA sequences. Substantially pure DNA also includes a recombinant DNA which is part of a hybrid gene encoding additional *S. epidermidis* DNA sequence.

A "contig" as used herein is a nucleic acid representing a continuous stretch of genomic sequence of an organism.

An "open reading frame", also referred to herein as ORF, is a region of nucleic acid which encodes a polypeptide. This region may represent a portion of a coding sequence or a total sequence and can be determined from a stop to stop codon or from a start to stop codon.

As used herein, a "coding sequence" is a nucleic acid which is transcribed into messenger RNA and/or translated into a polypeptide when placed under the control of appropriate regulatory sequences. The boundaries of the coding sequence are determined by a translation start codon at the five prime terminus and a translation stop code at the three prime terminus. A coding sequence can include but is not limited to messenger RNA, synthetic DNA, and recombinant nucleic acid sequences.

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A "complement" of a nucleic acid as used herein refers to an anti-parallel or antisense sequence that participates in Watson-Crick base-pairing with the original sequence.

A "gene product" is a protein or structural RNA which is specifically encoded by a gene.

As used herein, the term "probe" refers to a nucleic acid, peptide or other chemical entity which specifically binds to a molecule of interest. Probes are often associated with or capable of associating with a label. A label is a chemical moiety capable of detection. Typical labels comprise dyes, radioisotopes, luminescent and chemiluminescent moieties, fluorophores, enzymes, precipitating agents, amplification sequences, and the like. Similarly, a nucleic acid, peptide or other chemical entity which specifically binds to a molecule of interest and immobilizes such molecule is referred herein as a "capture ligand". Capture ligands are typically associated with or capable of associating with a support such as nitrocellulose, glass, nylon membranes, beads, particles and the like. The specificity of hybridization is dependent on conditions such as the base pair composition of the nucleotides, and the temperature and salt concentration of the reaction. These conditions are readily discernable to one of ordinary skill in the art using routine experimentation.

"Homologous" refers to the sequence similarity or sequence identity between two polypeptides or between two nucleic acid molecules. When a position in both of the two

compared sequences is occupied by the same base or amino acid monomer subunit, e.g., if a position in each of two DNA molecules is occupied by adenine, then the molecules are homologous at that position. The percent of homology between two sequences is a function of the number of matching or homologous positions shared by the two sequences divided by the number of positions compared x 100. For example, if 6 of 10 of the positions in two sequences are matched or homologous then the two sequences are 60% homologous. By way of example, the DNA sequences ATTGCC and TATGGC share 50% homology. Generally, a comparison is made when two sequences are aligned to give maximum homology.

Nucleic acids are hybridizable to each other when at least one strand of a nucleic acid

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can anneal to the other nucleic acid under defined stringency conditions. Stringency of hybridization is determined by: (a) the temperature at which hybridization and/or washing is performed; and (b) the ionic strength and polarity of the hybridization and washing solutions. Hybridization requires that the two nucleic acids contain complementary sequences; depending on the stringency of hybridization, however, mismatches may be tolerated.
Typically, hybridization of two sequences at high stringency (such as, for example, in a solution of 0.5X SSC, at 65° C) requires that the sequences be essentially completely homologous. Conditions of intermediate stringency (such as, for example, 2X SSC at 65° C) and low stringency (such as, for example 2X SSC at 55° C) require correspondingly less overall complementarity between the hybridizing sequences. (1X SSC is 0.15 M NaCl, 0.015
M Na citrate).

The terms peptides, proteins, and polypeptides are used interchangeably herein.

As used herein, the term "surface protein" refers to all surface accessible proteins, e.g. inner and outer membrane proteins, proteins adhering to the cell wall, and secreted proteins.

A polypeptide has *S. epidermidis* biological activity if it has one, two or preferably more of the following properties: (1) if when expressed in the course of an *S. epidermidis* infection, it can promote, or mediate the attachment of *S. epidermidis* to a cell; (2) it has an enzymatic activity, structural or regulatory function characteristic of an *S. epidermidis*

protein; (3) the gene which encodes it can rescue a lethal mutation in an *S. epidermidis* gene. A polypeptide has biological activity if it is an antagonist, agonist, or super-agonist of a polypeptide having one of the above-listed properties.

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A biologically active fragment or analog is one having an *in vivo* or *in vitro* activity which is characteristic of the *S. epidermidis* polypeptides of the invention contained in the Sequence Listing, or of other naturally occurring *S. epidermidis* polypeptides, e.g., one or more of the biological activities described herein. Especially preferred are fragments which exist *in vivo*, e.g., fragments which arise from post transcriptional processing or which arise from translation of alternatively spliced RNA's. Fragments include those expressed in native or endogenous cells as well as those made in expression systems, e.g., in CHO (Chinese Hamster Ovary) cells. Because peptides such as *S. epidermidis* polypeptides often exhibit a range of physiological properties and because such properties may be attributable to different portions of the molecule, a useful *S. epidermidis* fragment or *S. epidermidis* analog is one which exhibits a biological activity in any biological assay for *S. epidermidis* activity. The fragment or analog possesses about 10%, preferably about 40%, more preferably about 60%, 70%, 80% or 90% or greater of the activity of *S. epidermidis*, in any *in vivo* or *in vitro* assay.

Analogs can differ from naturally occurring *S. epidermidis* polypeptides in amino acid sequence or in ways that do not involve sequence, or both. Non-sequence modifications include changes in acetylation, methylation, phosphorylation, carboxylation, or glycosylation. Preferred analogs include *S. epidermidis* polypeptides (or biologically active fragments thereof) whose sequences differ from the wild-type sequence by one or more conservative amino acid substitutions or by one or more non-conservative amino acid substitutions, deletions, or insertions which do not substantially diminish the biological activity of the *S. epidermidis* polypeptide. Conservative substitutions typically include the substitution of one amino acid for another with similar characteristics, e.g., substitutions within the following groups: valine, glycine; glycine, alanine; valine, isoleucine, leucine; aspartic acid, glutamic

acid; asparagine, glutamine; serine, threonine; lysine, arginine; and phenylalanine, tyrosine.

Other conservative substitutions can be made in view of the table below.

TABLE 1
CONSERVATIVE AMINO ACID REPLACEMENTS

For Amino Acid	Code	Replace with any of
Alanine	A	D-Ala, Gly, beta-Ala, L-Cys, D-Cys
Arginine	R	D-Arg, Lys, D-Lys, homo-Arg, D-homo-Arg, Met, Ile, D-Met, D-Ile, Orn, D-Orn
Asparagine	N	D-Asn, Asp, D-Asp, Glu, D-Glu, Gln, D-Gln
Aspartic Acid	Q	D-Asp, D-Asn, Asn, Glu, D-Glu, Gln, D-Gln
Cysteine	С	D-Cys, S-Me-Cys, Met, D-Met, Thr, D-Thr
Glutamine	Q	D-Gln, Asn, D-Asn, Glu, D-Glu, Asp, D-Asp
Glutamic Acid	Е	D-Glu, D-Asp, Asp, Asn, D-Asn, Gln, D-Gln
Glycine	G	Ala, D-Ala, Pro, D-Pro, β-Ala, Acp
Isoleucine	I	D-Ile, Val, D-Val, Leu, D-Leu, Met, D-Met
Leucine	L	D-Leu, Val, D-Val, Leu, D-Leu, Met, D-Met
Lysine	K	D-Lys, Arg, D-Arg, homo-Arg, D-homo-Arg, Met, D-Met, Ile, D-Ile, Orn, D-Orn
Methionine	M	D-Met, S-Me-Cys, Ile, D-Ile, Leu, D-Leu, Val, D-Val
Phenylalanine	F	D-Phe, Tyr, D-Thr, L-Dopa, His, D-His, Trp, D-Trp, Trans-3,4, or 5-phenylproline, cis-3,4, or 5-phenylproline
Proline	P	D-Pro, L-I-thioazolidine-4-carboxylic acid, D-or L-1-oxazolidine-4-carboxylic acid
Serine	S	D-Ser, Thr, D-Thr, allo-Thr, Met, D-Met, Met(O), D-Met(O), L-Cys, D-Cys
Threonine	T	D-Thr, Ser, D-Ser, allo-Thr, Met, D-Met, Met(O), D-Met(O), Val, D-Val
Tyrosine	Y	D-Tyr, Phe, D-Phe, L-Dopa, His, D-His
Valine	V	D-Val, Leu, D-Leu, Ile, D-Ile, Met, D-Met

Other analogs within the invention are those with modifications which increase peptide stability; such analogs may contain, for example, one or more non-peptide bonds (which replace the peptide bonds) in the peptide sequence. Also included are: analogs that include residues other than naturally occurring L-amino acids, e.g., D-amino acids or non-naturally occurring or synthetic amino acids, e.g., β or γ amino acids; and cyclic analogs.

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As used herein, the term "fragment", as applied to an *S. epidermidis* analog, will ordinarily be at least about 20 residues, more typically at least about 40 residues, preferably at least about 60 residues in length. Fragments of *S. epidermidis* polypeptides can be generated by methods known to those skilled in the art. The ability of an *Staphylococcus* fragment to exhibit a biological activity of *S. epidermidis* polypeptide can be assessed by methods known to those skilled in the art as described herein. Also included are *S. epidermidis* polypeptides containing residues that are not required for biological activity of the peptide or that result from alternative mRNA splicing or alternative protein processing events.

An "immunogenic component" as used herein is a moiety, such as an *S. epidermidis* polypeptide, analog or fragment thereof, that is capable of eliciting a humoral and/or cellular immune response in a host animal.

An "antigenic component" as used herein is a moiety, such as an *S. epidermidis* polypeptide, analog or fragment thereof, that is capable of binding to a specific antibody with sufficiently high affinity to form a detectable antigen-antibody complex.

The term "antibody" as used herein is intended to include fragments thereof which are specifically reactive with *S. epidermidis* polypeptides.

As used herein, the term "cell-specific promoter" means a DNA sequence that serves as a promoter, i.e., regulates expression of a selected DNA sequence operably linked to the promoter, and which effects expression of the selected DNA sequence in specific cells of a tissue. The term also covers so-called "leaky" promoters, which regulate expression of a selected DNA primarily in one tissue, but cause expression in other tissues as well.

Misexpression, as used herein, refers to a non-wild type pattern of gene expression. It includes: expression at non-wild type levels, i.e., over or under expression; a pattern of expression that differs from wild type in terms of the time or stage at which the gene is expressed, e.g., increased or decreased expression (as compared with wild type) at a predetermined developmental period or stage; a pattern of expression that differs from wild type in terms of increased expression (as compared with wild type) in a predetermined cell type or tissue type; a pattern of expression that differs from wild type in terms of the splicing size, amino acid sequence, post-translational modification, or biological activity of the expressed polypeptide; a pattern of expression that differs from wild type in terms of the effect of an environmental stimulus or extracellular stimulus on expression of the gene, e.g., a pattern of increased or decreased expression (as compared with wild type) in the presence of an increase or decrease in the strength of the stimulus.

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As used herein, "host cells" and other such terms denoting microorganisms or higher eukaryotic cell lines cultured as unicellular entities refers to cells which can become or have been used as recipients for a recombinant vector or other transfer DNA, and include the progeny of the original cell which has been transfected. It is understood by individuals skilled in the art that the progeny of a single parental cell may not necessarily be completely identical in genomic or total DNA compliment to the original parent, due to accident or deliberate mutation.

As used herein, the term "control sequence" refers to a nucleic acid having a base sequence which is recognized by the host organism to effect the expression of encoded sequences to which they are ligated. The nature of such control sequences differs depending upon the host organism; in prokaryotes, such control sequences generally include a promoter, ribosomal binding site, terminators, and in some cases operators; in eukaryotes, generally such control sequences include promoters, terminators and in some instances, enhancers. The term control sequence is intended to include at a minimum, all components whose presence is

necessary for expression, and may also include additional components whose presence is advantageous, for example, leader sequences.

As used herein, the term "operably linked" refers to sequences joined or ligated to function in their intended manner. For example, a control sequence is operably linked to coding sequence by ligation in such a way that expression of the coding sequence is achieved under conditions compatible with the control sequence and host cell.

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The "metabolism" of a substance, as used herein, means any aspect of the expression, function, action, or regulation of the substance. The metabolism of a substance includes modifications, e.g., covalent or non-covalent modifications of the substance. The metabolism of a substance includes modifications, e.g., covalent or non-covalent modification, the substance induces in other substances. The metabolism of a substance also includes changes in the distribution of the substance. The metabolism of a substance includes changes the substance induces in the distribution of other substances.

A "sample" as used herein refers to a biological sample, such as, for example, tissue or fluid isloated from an individual (including without limitation plasma, serum, cerebrospinal fluid, lymph, tears, saliva and tissue sections) or from *in vitro* cell culture constituents, as well as samples from the environment.

Technical and scientific terms used herein have the meanings commonly understood by one of ordinary skill in the art to which the present invention pertains, unless otherwise defined. Reference is made herein to various methodologies known to those of skill in the art. Publications and other materials setting forth such known methodologies to which reference is made are incorporated herein by reference in their entireties as though set forth in full. The practice of the invention will employ, unless otherwise indicated, conventional techniques of chemistry, molecular biology, microbiology, recombinant DNA, and immunology, which are within the skill of the art. Such techniques are explained fully in the literature. *See e.g.*, Sambrook, Fritsch, and Maniatis, *Molecular Cloning; Laboratory Manual* 2nd ed. (1989); *DNA Cloning*, Volumes I and II (D.N Glover ed. 1985); *Oligonucleotide Synthesis* (M.J. Gait

ed, 1984); Nucleic Acid Hybridization (B.D. Hames & S.J. Higgins eds. 1984); the series, Methods in Enzymology (Academic Press, Inc.), particularly Vol. 154 and Vol. 155 (Wu and Grossman, eds.); PCR-A Practical Approach (McPherson, Quirke, and Taylor, eds., 1991); Immunology, 2d Edition, 1989, Roitt et al., C.V. Mosby Company, and New York; Advanced 5 Immunology, 2d Edition, 1991, Male et al., Grower Medical Publishing, New York.: DNA Cloning: A Practical Approach, Volumes I and II, 1985 (D.N. Glover ed.); Oligonucleotide Synthesis, 1984, (M.L. Gait ed); Transcription and Translation, 1984 (Hames and Higgins eds.); Animal Cell Culture, 1986 (R.I. Freshney ed.); Immobilized Cells and Enzymes, 1986 (IRL Press); Perbal, 1984, A Practical Guide to Molecular Cloning; Gene Transfer Vectors 10 for Mammalian Cells, 1987 (J. H. Miller and M. P. Calos eds., Cold Spring Harbor Laboratory); Martin J. Bishop, ed., Guide to Human Genome Computing, 2d Edition, Academic Press, San Diego, CA. (1998); and Leonard F. Peruski, Jr., and Anne Harwood Peruski, The Internet and the New Biology: Tools for Genomic and Molecular Research, American Society for Microbiology, Washington, D.C. (1997).

Any suitable materials and/or methods known to those of skill can be utilized in carrying out the present invention; however, preferred materials and/or methods are described.

Materials, reagents and the like to which reference is made in the following description and examples are obtainable from commercial sources, unless otherwise noted.

20 S. epidermidis Genomic Sequence

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This invention provides nucleotide sequences of the genome of *S. epidermidis* which thus comprises a DNA sequence library of *S. epidermidis* genomic DNA. The detailed description that follows provides nucleotide sequences of *S. epidermidis*, and also describes how the sequences were obtained and how ORFs and protein-coding sequences were identified. Also described are compositions and methods of using the disclosed *S. epidermidis* sequences in methods including diagnostic and therapeutic applications.

Furthermore, the library can be used as a database for identification and comparison of medically important sequences in this and other strains of *S. epidermidis*.

To determine the genomic sequence of *S. epidermidis*, DNA from strain 18972 of *S. epidermidis* was isolated and a library of DNA fragments were transformed into DH5α cells. DNA sequencing was achieved using established ABI sequencing methods on ABI377 automated DNA sequencers. The cloning and sequencing procedures are described in more detail in the Exemplification.

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Individual sequence reads were assembled using PHRAP (P. Green, Abstracts of DOE Human Genome Program Contractor-Grantee Workshop V, Jan. 1996, p.157). The average contig length was about 3-4 kb.

All subsequent steps were based on sequencing by ABI377 automated DNA sequencing methods. The cloning and sequencing procedures are described in more detail in the Exemplification.

A variety of approaches may be used to order the contigs so as to obtain a continuous sequence representing the entire *S. epidermidis* genome. Synthetic oligonucleotides are designed that are complementary to sequences at the end of each contig. These oligonucleotides may be hybridized to libaries of *S. epidermidis* genomic DNA in, for example, lambda phage vectors or plasmid vectors to identify clones that contain sequences corresponding to the junctional regions between individual contigs. Such clones are then used to isolate template DNA and the same oligonucleotides are used as primers in polymerase chain reaction (PCR) to amplify junctional fragments, the nucleotide sequence of which is then determined.

The *S. epidermidis* sequences were analyzed for the presence of open reading frames (ORFs) comprising at least 180 nucleotides. As a result of the analysis of ORFs based on stop-to-stop codon reads, it should be understood that these ORFs may not correspond to the ORF of a naturally-occurring *S. epidermidis* polypeptide. These ORFs may contain start codons which indicate the initiation of protein synthesis of a naturally-occurring *S.*

epidermidis polypeptide. Such start codons within the ORFs provided herein were identified by those of ordinary skill in the relevant art, and the resulting ORF and the encoded *S. epidermidis* polypeptide is within the scope of this invention. For example, within the ORFs a codon such as AUG or GUG (encoding methionine or valine) which is part of the initiation signal for protein synthesis were identified and the portion of an ORF to corresponding to a naturally-occurring *S. epidermidis* polypeptide was recognized. The predicted coding regions were defined by evaluating the coding potential of such sequences with the program GENEMARKTM (Borodovsky and McIninch, 1993, *Comp.* . <u>17</u>:123).

Each predicted ORF amino acid sequence was compared with all sequences found in current GENBANK, SWISS-PROT, and PIR databases using the BLAST algorithm. BLAST identifies local alignments occurring by chance between the ORF sequence and the sequence in the databank (Altschal et al., 1990, L Mol. Biol. 215:403-410). Homologous ORFs (probabilities less than 10⁻⁵ by chance) and ORF's that are probably non-homologous (probabilities greater than 10⁻⁵ by chance) but have good codon usage were identified. Both homologous, sequences and non-homologous sequences with good codon usage, are likely to encode proteins and are encompassed by the invention.

S. epidermidis Plasmid Sequences

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This invention also provides nucleotide sequences of two naturally occurs plasmids

which thus comprises a DNA sequence library of *S. epidermidis* plasmid DNA. One plasmid disclosed is approximately 39 Kb and nucleic acid sequence is contained within SEQ ID NO: 3703 - SEQ ID NO: 3764. The other plasmid is approximately 2.9 Kb contained within SEQ ID NO: 3765 - SEQ ID NO: 3772. The detailed description that follows provides nucleotide sequences of *S. epidermidis*, and also describes how the sequences were obtained and how ORFs and protein-coding sequences were identified. Also described are compositions and methods of using the disclosed *S. epidermidis* sequences in methods including diagnostic and therapeutic applications. The plasmid sequences can also be used as vectors and gene

expression. Furthermore, the plasmid library can be used as a database for identification and comparison of medically important sequences in this and other strains of *S. epidermidis*.

Similar methods were used to determine to plasmid sequences of *S. epidermidis* as described above in determining the genomic sequence. A more detailed description of the methods are in the Exemplification.

S. epidermidis Nucleic Acids

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The present invention provides a library of *S. epidermidis* -derived nucleic acid sequences. The libraries provide probes, primers, and markers which are used as markers in epidemiological studies. The present invention also provides a library of *S. epidermidis* - derived nucleic acid sequences which comprise or encode targets for therapeutic drugs.

The nucleic acids of this invention may be obtained directly from the DNA of the above referenced *S. epidermidis* strain by using the polymerase chain reaction (PCR). See "PCR, A Practical Approach" (McPherson, Quirke, and Taylor, eds., IRL Press, Oxford, UK, 1991) for details about the PCR. High fidelity PCRis used to ensure a faithful DNA copy prior to expression. In addition, the authenticity of amplified products is verified by conventional sequencing methods. Clones carrying the desired sequences described in this invention may also be obtained by screening the libraries by means of the PCR or by hybridization of synthetic oligonucleotide probes to filter lifts of the library colonies or plaques as known in the art (see, e.g., Sambrook et al., Molecular Cloning, A Laboratory Manual 2nd edition, 1989, Cold Spring Harbor Press, NY).

It is also possible to obtain nucleic acids encoding *S. epidermidis* polypeptides from a cDNA library in accordance with protocols herein described. A cDNA encoding an *S. epidermidis* polypeptide can be obtained by isolating total mRNA from an appropriate strain.

Double stranded cDNAs can then be prepared from the total mRNA. Subsequently, the cDNAs can be inserted into a suitable plasmid or viral (e.g., bacteriophage) vector using any one of a number of known techniques. Genes encoding *S. epidermidis* polypeptides can also

be cloned using established polymerase chain reaction techniques in accordance with the nucleotide sequence information provided by the invention. The nucleic acids of the invention can be DNA or RNA. Preferred nucleic acids of the invention are contained in the Sequence Listing.

The nucleic acids of the invention can also be chemically synthesized using standard techniques. Various methods of chemically synthesizing polydeoxynucleotides are known, including solid-phase synthesis which, like peptide synthesis, has been fully automated in commercially available DNA synthesizers (See e.g., Itakura et al. U.S. Patent No. 4,598,049; Caruthers et al. U.S. Patent No. 4,458,066; and Itakura U.S. Patent Nos. 4,401,796 and 4,373,071, incorporated by reference herein).

In another example, DNA can be chemically synthesized using, e.g., the phosphoramidite solid support method of Matteucci *et al.*, 1981, *J. Am. Chem. Soc.* 103:3185, the method of Yoo *et al.*, 1989, *J. Biol. Chem.* 764:17078, or other well known methods. This can be done by sequentially linking a series of oligonucleotide cassettes comprising pairs of synthetic oligonucleotides, as described below.

Nucleic acids isolated or synthesized in accordance with features of the present invention are useful, by way of example, without limitation, as probes, primers, capture ligands, antisense genes and for developing expression systems for the synthesis of proteins and peptides corresponding to such sequences. As probes, primers, capture ligands and antisense agents, the nucleic acid normally consists of all or part (approximately twenty or more nucleotides for specificity as well as the ability to form stable hybridization products) of the nucleic acids of the invention contained in the Sequence Listing. These uses are described in further detail below.

25 Probes

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A nucleic acid isolated or synthesized in accordance with the sequence of the invention contained in the Sequence Listing can be used as a probe to specifically detect S.

epidermidis. With the sequence information set forth in the present application, sequences of twenty or more nucleotides are identified which provide the desired inclusivity and exclusivity with respect to *S. epidermidis*, and extraneous nucleic acids likely to be encountered during hybridization conditions. More preferably, the sequence will comprise at least about twenty to thirty nucleotides to convey stability to the hybridization product formed between the probe and the intended target molecules.

Sequences larger than 1000 nucleotides in length are difficult to synthesize but can be generated by recombinant DNA techniques. Individuals skilled in the art will readily recognize that the nucleic acids, for use as probes, can be provided with a label to facilitate detection of a hybridization product.

Nucleic acid isolated and synthesized in accordance with the sequence of the invention contained in the Sequence Listing can also be useful as probes to detect homologous regions (especially homologous genes) of other *Staphylococcus* species using appropriate stringency hybridization conditions as described herein.

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Capture Ligand

For use as a capture ligand, the nucleic acid selected in the manner described above with respect to probes, can be readily associated with a support. The manner in which nucleic acid is associated with supports is well known. Nucleic acid having twenty or more nucleotides in a sequence of the invention contained in the Sequence Listing have utility to separate *S. epidermidis* nucleic acid from one strain from the nucleic acid of other another strain as well as from other organisms. Nucleic acid having twenty or more nucleotides in a sequence of the invention contained in the Sequence Listing can also have utility to separate other *Staphylococcus* species from each other and from other organisms. Preferably, the sequence will comprise at least about twenty nucleotides to convey stability to the hybridization product formed between the probe and the intended target molecules.

Sequences larger than 1000 nucleotides in length are difficult to synthesize but can be generated by recombinant DNA techniques.

Primers

Nucleic acid isolated or synthesized in accordance with the sequences described herein have utility as primers for the amplification of *S. epidermidis* nucleic acid. These nucleic acids may also have utility as primers for the amplification of nucleic acids in other *Staphylococcus* species. With respect to polymerase chain reaction (PCR) techniques, nucleic acid sequences of ≥ 10-15 nucleotides of the invention contained in the Sequence

10 Listing have utility in conjunction with suitable enzymes and reagents to create copies of *S. epidermidis* nucleic acid. More preferably, the sequence will comprise twenty or more nucleotides to convey stability to the hybridization product formed between the primer and the intended target molecules. Binding conditions of primers greater than 100 nucleotides are more difficult to control to obtain specificity. High fidelity PCR can be used to ensure a faithful DNA copy prior to expression. In addition, amplified products can be checked by conventional sequencing methods.

The copies can be used in diagnostic assays to detect specific sequences, including genes from *S. epidermidis* and/or other *Staphylococcus* species. The copies can also be incorporated into cloning and expression vectors to generate polypeptides corresponding to the nucleic acid synthesized by PCR, as is described in greater detail herein.

The nucleic acids of the present invention find use as templates for the recombinant production of *S. epidermidis* -derived peptides or polypeptides

Antisense

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Nucleic acid or nucleic acid-hybridizing derivatives isolated or synthesized in accordance with the sequences described herein have utility as antisense agents to prevent the

expression of *S. epidermidis* genes. These sequences also have utility as antisense agents to prevent expression of genes of other *Staphylococcus* species.

In one embodiment, nucleic acid or derivatives corresponding to *S. epidermidis* nucleic acids is loaded into a suitable carrier such as a liposome or bacteriophage for introduction into bacterial cells. For example, a nucleic acid having twenty or more nucleotides is capable of binding to bacteria nucleic acid or bacteria messenger RNA. Preferably, the antisense nucleic acid is comprised of 20 or more nucleotides to provide necessary stability of a hybridization product of non-naturally occurring nucleic acid and bacterial nucleic acid and/or bacterial messenger RNA. Nucleic acid having a sequence greater than 1000 nucleotides in length is difficult to synthesize but can be generated by recombinant DNA techniques. Methods for loading antisense nucleic acid in liposomes is known in the art as exemplified by U.S. Patent 4,241,046 issued December 23, 1980 to Papahadjopoulos et al.

The present invention encompasses isolated polypeptides and nucleic acids derived from *S. epidermidis* that are useful as reagents for diagnosis of bacterial infection, components of effective anti-bacterial vaccines, and/or as targets for anti-bacterial drugs, including anti-*S. epidermidis* drugs.

Expression of S. epidermidis Nucleic Acids

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Table 2, which is appended herewith and which forms part of the present specification, provides a list of open reading frames (ORFs) in both strands and a putative identification of the particular function of a polypeptide which is encoded by each ORF, based on the homology match (determined by the BLASTP2 algorithm) of the predicted polypeptide with known proteins encoded by ORFs in other organisms. An ORF is a region of nucleic acid which encodes a polypeptide. This region may represent a portion of a coding sequence or a total sequence and was determined from stop to stop codons. The first column contains a

designation for the ORF ("ORF Name"). The second and third columns list the SEQ ID numbers for the nucleic acid ("NT ID") and amino acid ("AA ID") sequences corresponding to each ORF, respectively. The fourth and fifth columns list the length of the nucleic acid ORF ("NT LN") and the length of the amino acid ORF ("AA LN"), respectively. The 5 nucleotide sequence corresponding to each ORF begins at the first nucleotide immediately following a stop codon and ends at the nucleotide immediately preceding the next downstream stop codon in the same reading frame. It will be recognized by one skilled in the art that the natural translation initiation sites will correspond to ATG, GTG, or TTG codons located within the ORFs. The natural initiation sites depend not only on the sequence of a start codon 10 but also on the context of the DNA sequence adjacent to the start codon. Usually, a recognizable ribosome binding site is found within 20 nucleotides upstream from the initiation codon. In some cases where genes are translationally coupled and coordinately expressed together in "operons", ribosome binding sites are not present, but the initiation codon of a downstream gene may occur very close to, or overlap, the stop codon of the an upstream gene in the same operon. The correct start codons can be generally identified without undue 15 experimentation because only a few codons need be tested. It is recognized that the translational machinery in bacteria initiates all polypeptide chains with the amino acid methionine, regardless of the sequence of the start codon. In some cases, polypeptides are post-translationally modified, resulting in an N-terminal amino acid other than methionine in 20 vivo. The sixth and seventh columns provide metrics for assessing the likelihood of the homology match (determined by the BLASTP2 algorithm), as is known in the art, to the genes indicated in the description frame ("Description") below the ORF Name. These genes in the Description were identified when the designated ORF was compared against a comprehensive

non-redundant protein database. Specifically, the sixth column represents the Blast Score ("Score") for the match (a higher score is a better match), and the seventh column represents the probability ("P-value") for the match (the probability that such a match can have occurred by chance; the lower the value, the more likely the match is valid). If a BLASTP2 score of less than 100 was obtained, no value is reported in the table. The Description provides, where available, the Swissprot accession number (SP), the locus name (LN), the Organism (OR), Source of variant (SR), E.C. number (EC), the gene name (GN), the product name (PN), the Function Description (FN), Left End (LE), Right End (RE), Coding Direction (DI), and the description (DE) or notes (NT) for each ORF. This information allows one of ordinary skill in the art to determine a potential use for each identified coding sequence and, as a result, allows to use the polypeptides of the present invention for commercial and industrial purposes.

Using the information provided in SEQ ID NO: 1 - SEQ ID NO: 3772, SEQ ID NO: 3773 - SEQ ID NO: 7544 and in Table 2 together with routine cloning and sequencing methods, one of ordinary skill in the art will be able to clone and sequence all the nucleic acid fragments of interest including open reading frames (ORFs) encoding a large variety of proteins of *S. epidermidis*.

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Nucleic acid isolated or synthesized in accordance with the sequences described herein have utility to generate polypeptides. The nucleic acid of the invention exemplified in SEQ ID NO: 1 - SEQ ID NO: 3772 and in Table 2 or fragments of said nucleic acid encoding active portions of *S. epidermidis* polypeptides can be cloned into suitable vectors or used to isolate nucleic acid. The isolated nucleic acid is combined with suitable DNA linkers and cloned into a suitable vector.

The function of a specific gene or operon can be ascertained by expression in a bacterial strain under conditions where the activity of the gene product(s) specified by the gene or operon in question can be specifically measured. Alternatively, a gene product may be produced in large quantities in an expressing strain for use as an antigen, an industrial reagent, for structural studies, etc. This expression can be accomplished in a mutant strain which lacks the activity of the gene to be tested, or in a strain that does not produce the same gene product(s). This includes, but is not limited to, Eucaryotic species such as the yeast *Saccharomyces cerevisiae*, *Methanobacterium* strains or other Archaea, and Eubacteria such as *E. coli*, *B. Subtilis*, *S. Aureus*, *S. Pneumonia* or *Pseudomonas putida*. In some cases the expression host will utilize the natural *S. epidermidis* promoter whereas in others, it will be necessary to drive the gene with a promoter sequence derived from the expressing organism (e.g., an *E. coli* beta-galactosidase promoter for expression in *E. coli*).

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To express a gene product using the natural *S. epidermidis* promoter, a procedure such as the following can be used. A restriction fragment containing the gene of interest, together with its associated natural promoter element and regulatory sequences (identified using the DNA sequence data) is cloned into an appropriate recombinant plasmid containing an origin of replication that functions in the host organism and an appropriate selectable marker. This can be accomplished by a number of procedures known to those skilled in the art. It is most preferably done by cutting the plasmid and the fragment to be cloned with the same restriction enzyme to produce compatible ends that can be ligated to join the two pieces together. The recombinant plasmid is introduced into the host organism by, for example, electroporation and cells containing the recombinant plasmid are identified by selection for the marker on the plasmid. Expression of the desired gene product is detected using an assay specific for that gene product.

In the case of a gene that requires a different promoter, the body of the gene (coding sequence) is specifically excised and cloned into an appropriate expression plasmid. This subcloning can be done by several methods, but is most easily accomplished by PCR

amplification of a specific fragment and ligation into an expression plasmid after treating the PCR product with a restriction enzyme or exonuclease to create suitable ends for cloning.

A suitable host cell for expression of a gene can be any procaryotic or eucaryotic cell. Suitable methods for transforming host cells can be found in Sambrook et al. (Molecular Cloning: A Laboratory Manual, 2nd Edition, Cold Spring Harbor Laboratory Press (1989)), and other laboratory textbooks.

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For example, a host cell transfected with a nucleic acid vector directing expression of a nucleotide sequence encoding an *S. epidermidis* polypeptide can be cultured under appropriate conditions to allow expression of the polypeptide to occur. Suitable media for cell culture are well known in the art. Polypeptides of the invention can be isolated from cell culture medium, host cells, or both using techniques known in the art for purifying proteins including ion-exchange chromatography, gel filtration chromatography, ultrafiltration, electrophoresis, and immunoaffinity purification with antibodies specific for such polypeptides. Additionally, in many situations, polypeptides can be produced by chemical cleavage of a native protein (e.g., tryptic digestion) and the cleavage products can then be purified by standard techniques.

In the case of membrane bound proteins, these can be isolated from a host cell by contacting a membrane-associated protein fraction with a detergent forming a solubilized complex, where the membrane-associated protein is no longer entirely embedded in the membrane fraction and is solubilized at least to an extent which allows it to be chromatographically isolated from the membrane fraction. Chromatographic techniques which can be used in the final purification step are known in the art and include hydrophobic interaction, lectin affinity, ion exchange, dye affinity and immunoaffinity.

One strategy to maximize recombinant *S. epidermidis* peptide expression in *E. coli* is to express the protein in a host bacteria with an impaired capacity to proteolytically cleave the recombinant protein (Gottesman, S., <u>Gene Expression Technology: Methods in Enzymology 185</u>, Academic Press, San Diego, California (1990) 119-128). Another strategy would be to

alter the nucleic acid encoding an *S. epidermidis* peptide to be inserted into an expression vector so that the individual codons for each amino acid would be those preferentially utilized in highly expressed *E. coli* proteins (Wada et al., (1992) *Nuc. Acids Res.* 20:2111-2118). Such alteration of nucleic acids of the invention can be carried out by standard DNA synthesis techniques.

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The nucleic acids of the invention can also be chemically synthesized using standard techniques. Various methods of chemically synthesizing polydeoxynucleotides are known, including solid-phase synthesis which, like peptide synthesis, has been fully automated in commercially available DNA synthesizers (See, e.g., Itakura et al. U.S. Patent No. 4,598,049; Caruthers et al. U.S. Patent No. 4,458,066; and Itakura U.S. Patent Nos. 4,401,796 and 4,373,071, incorporated by reference herein).

The present invention provides a library of *S. epidermidis* -derived nucleic acid sequences. The libraries provide probes, primers, and markers which can be used as markers in epidemiological studies. The present invention also provides a library of *S. epidermidis* - derived nucleic acid sequences which comprise or encode targets for therapeutic drugs.

Nucleic acids comprising any of the sequences disclosed herein or sub-sequences thereof can be prepared by standard methods using the nucleic acid sequence information provided in SEQ ID NO: 1 - SEQ ID NO: 3772. For example, DNA can be chemically synthesized using, e.g., the phosphoramidite solid support method of Matteucci *et al.*, 1981, *J. Am. Chem. Soc.* 103:3185, the method of Yoo *et al.*, 1989, *J. Biol. Chem.* 764:17078, or other well known methods. This can be done by sequentially linking a series of oligonucleotide cassettes comprising pairs of synthetic oligonucleotides, as described below.

Of course, due to the degeneracy of the genetic code, many different nucleotide sequences can encode polypeptides having the amino acid sequences defined by SEQ ID NO: 3773 - SEQ ID NO: 7544 or sub-sequences thereof. The codons can be selected for optimal expression in prokaryotic or eukaryotic systems. Such degenerate variants are also encompassed by this invention.

Insertion of nucleic acids (typically DNAs) encoding the polypeptides of the invention into a vector is easily accomplished when the termini of both the DNAs and the vector comprise compatible restriction sites. If this cannot be done, it may be necessary to modify the termini of the DNAs and/or vector by digesting back single-stranded DNA overhangs generated by restriction endonuclease cleavage to produce blunt ends, or to achieve the same result by filling in the single-stranded termini with an appropriate DNA polymerase.

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Alternatively, any site desired may be produced, e.g., by ligating nucleotide sequences (linkers) onto the termini. Such linkers may comprise specific oligonucleotide sequences that define desired restriction sites. Restriction sites can also be generated by the use of the polymerase chain reaction (PCR). See, e.g., Saiki *et al.*, 1988, *Science* 239:48. The cleaved vector and the DNA fragments may also be modified if required by homopolymeric tailing.

The nucleic acids of the invention may be isolated directly from cells. Alternatively, the polymerase chain reaction (PCR) method can be used to produce the nucleic acids of the invention, using either chemically synthesized strands or genomic material as templates.

Primers used for PCR can be synthesized using the sequence information provided herein and can further be designed to introduce appropriate new restriction sites, if desirable, to facilitate incorporation into a given vector for recombinant expression.

The nucleic acids of the present invention may be flanked by natural *S. epidermidis* regulatory sequences, or may be associated with heterologous sequences, including promoters, enhancers, response elements, signal sequences, polyadenylation sequences, introns, 5'- and 3'- noncoding regions, and the like. The nucleic acids may also be modified by many means known in the art. Non-limiting examples of such modifications include methylation, "caps", substitution of one or more of the naturally occurring nucleotides with an analog, internucleotide modifications such as, for example, those with uncharged linkages (e.g., methyl phosphonates, phosphotriesters, phosphoroamidates, carbamates, etc.) and with charged linkages (e.g., phosphorothioates, phosphorodithioates, etc.). Nucleic acids may contain one or more additional covalently linked moieties, such as, for example, proteins (e.g.,

nucleases, toxins, antibodies, signal peptides, poly-L-lysine, etc.), intercalators (e.g., acridine, psoralen, etc.), chelators (e.g., metals, radioactive metals, iron, oxidative metals, etc.), and alkylators. PNAs are also included. The nucleic acid may be derivatized by formation of a methyl or ethyl phosphotriester or an alkyl phosphoramidate linkage. Furthermore, the nucleic acid sequences of the present invention may also be modified with a label capable of providing a detectable signal, either directly or indirectly. Exemplary labels include radioisotopes, fluorescent molecules, biotin, and the like.

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The invention also provides nucleic acid vectors comprising the disclosed *S. epidermidis* -derived sequences or derivatives or fragments thereof. A large number of vectors, including plasmid and bacterial vectors, have been described for replication and/or expression in a variety of eukaryotic and prokaryotic hosts, and may be used for cloning or protein expression.

The encoded *S. epidermidis* polypeptides may be expressed by using many known vectors, such as pUC plasmids, pET plasmids (Novagen, Inc., Madison, WI), or pRSET or pREP (Invitrogen, San Diego, CA), and many appropriate host cells, using methods disclosed or cited herein or otherwise known to those skilled in the relevant art. The particular choice of vector/host is not critical to the practice of the invention.

Recombinant cloning vectors will often include one or more replication systems for cloning or expression, one or more markers for selection in the host, e.g. antibiotic resistance, and one or more expression cassettes. The inserted *S. epidermidis* coding sequences may be synthesized by standard methods, isolated from natural sources, or prepared as hybrids, etc. Ligation of the *S. epidermidis* coding sequences to transcriptional regulatory elements and/or to other amino acid coding sequences may be achieved by known methods. Suitable host cells may be transformed/transfected/infected as appropriate by any suitable method including electroporation, CaCl₂ mediated DNA uptake, bacterial infection, microinjection, microprojectile, or other established methods.

Appropriate host cells include bacteria, archebacteria, fungi, especially yeast, and plant and animal cells, especially mammalian cells. Of particular interest are *S. epidermidis*, *E. coli, B. Subtilis, Saccharomyces cerevisiae, Saccharomyces carlsbergensis, Schizosaccharomyces pombi, SF9* cells, C129 cells, 293 cells, *Neurospora*, and CHO cells, COS cells, HeLa cells, and immortalized mammalian myeloid and lymphoid cell lines. Preferred replication systems include M13, ColE1, SV40, baculovirus, lambda, adenovirus, and the like. A large number of transcription initiation and termination regulatory regions have been isolated and shown to be effective in the transcription and translation of heterologous proteins in the various hosts. Examples of these regions, methods of isolation, manner of manipulation, etc. are known in the art. Under appropriate expression conditions, host cells can be used as a source of recombinantly produced *S. epidermidis* -derived peptides and polypeptides.

Advantageously, vectors may also include a transcription regulatory element (i.e., a promoter) operably linked to the *S. epidermidis* portion. The promoter may optionally contain operator portions and/or ribosome binding sites. Non-limiting examples of bacterial promoters compatible with *E. coli* include: b-lactamase (penicillinase) promoter; lactose promoter; tryptophan (trp) promoter; araBAD (arabinose) operon promoter; lambda-derived P₁ promoter and N gene ribosome binding site; and the hybrid tac promoter derived from sequences of the trp and lac UV5 promoters. Non-limiting examples of yeast promoters include 3-phosphoglycerate kinase promoter, glyceraldehyde-3-phosphate dehydrogenase (GAPDH) promoter, galactokinase (GAL1) promoter, galactoepimerase promoter, and alcohol dehydrogenase (ADH) promoter. Suitable promoters for mammalian cells include without limitation viral promoters such as that from Simian Virus 40 (SV40), Rous sarcoma virus (RSV), adenovirus (ADV), and bovine papilloma virus (BPV). Mammalian cells may also require terminator sequences, polyA addition sequences and enhancer sequences to increase expression. Sequences which cause amplification of the gene may also be desirable.

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including, but not limited to, bacteria, yeast, and animal cells, such as secretory signal sequences and/or prohormone pro region sequences, may also be included. These sequences are well described in the art.

Nucleic acids encoding wild-type or variant *S. epidermidis* -derived polypeptides may also be introduced into cells by recombination events. For example, such a sequence can be introduced into a cell, and thereby effect homologous recombination at the site of an endogenous gene or a sequence with substantial identity to the gene. Other recombination-based methods such as nonhomologous recombinations or deletion of endogenous genes by homologous recombination may also be used.

The nucleic acids of the present invention find use as templates for the recombinant production of *S. epidermidis* -derived peptides or polypeptides.

Identification and Use of S. epidermidis Nucleic Acid Sequences

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The disclosed *S. epidermidis* polypeptide and nucleic acid sequences, or other sequences that are contained within ORFs, including complete protein-coding sequences, of which any of the disclosed *S. epidermidis* -specific sequences forms a part, are useful as target components for diagnosis and/or treatment of *S. epidermidis* - caused infection

It will be understood that the sequence of an entire protein-coding sequence of which each disclosed nucleic acid sequence forms a part can be isolated and identified based on each disclosed sequence. This can be achieved, for example, by using an isolated nucleic acid encoding the disclosed sequence, or fragments thereof, to prime a sequencing reaction with genomic *S. epidermidis* DNA as template; this is followed by sequencing the amplified product. The isolated nucleic acid encoding the disclosed sequence, or fragments thereof, can also be hybridized to *S. epidermidis* genomic libraries to identify clones containing additional complete segments of the protein-coding sequence of which the shorter sequence forms a part. Then, the entire protein-coding sequence, or fragments thereof, or nucleic acids encoding all

or part of the sequence, or sequence-conservative or function-conservative variants thereof, may be employed in practicing the present invention.

Preferred sequences are those that are useful in diagnostic and/or therapeutic applications. Diagnostic applications include without limitation nucleic-acid-based and antibody-based methods for detecting bacterial infection. Therapeutic applications include without limitation vaccines, passive immunotherapy, and drug treatments directed against gene products that are both unique to bacteria and essential for growth and/or replication of bacteria.

10 Identification of Nucleic Acids Encoding Vaccine Components and Targets for Agents Effective Against S. epidermidis

The disclosed *S. epidermidis* genome sequence includes segments that direct the synthesis of ribonucleic acids and polypeptides, as well as origins of replication, promoters, other types of regulatory sequences, and intergenic nucleic acids. The invention encompasses nucleic acids encoding immunogenic components of vaccines and targets for agents effective against *S. epidermidis*. Identification of said immunogenic components involved in the determination of the function of the disclosed sequences, which can be achieved using a variety of approaches. Non-limiting examples of these approaches are described briefly below.

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Homology to known sequences:

Computer-assisted comparison of the disclosed *S. epidermidis* sequences with previously reported sequences present in publicly available databases is useful for identifying functional *S. epidermidis* nucleic acid and polypeptide sequences. It will be understood that protein-coding sequences, for example, may be compared as a whole, and that a high degree of sequence homology between two proteins (such as, for example, >80-90%) at the amino acid level indicates that the two proteins also possess some degree of functional homology,

such as, for example, among enzymes involved in metabolism, DNA synthesis, or cell wall synthesis, and proteins involved in transport, cell division, etc. In addition, many structural features of particular protein classes have been identified and correlate with specific consensus sequences, such as, for example, binding domains for nucleotides, DNA, metal ions, and other small molecules; sites for covalent modifications such as phosphorylation, acylation, and the like; sites of protein:protein interactions, etc. These consensus sequences may be quite short and thus may represent only a fraction of the entire protein-coding sequence. Identification of such a feature in an *S. epidermidis* sequence is therefore useful in determining the function of the encoded protein and identifying useful targets of antibacterial drugs.

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Of particular relevance to the present invention are structural features that are common to secretory, transmembrane, and surface proteins, including secretion signal peptides and hydrophobic transmembrane domains. *S. epidermidis* proteins identified as containing putative signal sequences and/or transmembrane domains are useful as immunogenic components of vaccines.

Targets for therapeutic drugs according to the invention include, but are not limited to, polypeptides of the invention, whether unique to *S. epidermidis* or not, that are essential for growth and/or viability of *S. epidermidis* under at least one growth condition. Polypeptides essential for growth and/or viability can be determined by examining the effect of deleting and/or disrupting the genes, i.e., by so-called gene "knockout". Alternatively, genetic footprinting can be used (Smith *et al.*, 1995, *Proc. Natl. Acad. Sci. USA* 92:5479-6433; Published International Application WO 94/26933; U.S. Patent No. 5,612,180). Still other methods for assessing essentiality includes the ability to isolate conditional lethal mutations in the specific gene (e.g., temperature sensitive mutations). Other useful targets for therapeutic drugs, which include polypeptides that are not essential for growth or viability *per se* but lead to loss of viability of the cell, can be used to target therapeutic agents to cells.

Strain-specific sequences:

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Because of the evolutionary relationship between different *S. epidermidis* strains, it is believed that the presently disclosed *S. epidermidis* sequences are useful for identifying, and/or discriminating between, previously known and new *S. epidermidis* strains. It is believed that other *S. epidermidis* strains will exhibit at least about 70% sequence homology with the presently disclosed sequence. Systematic and routine analyses of DNA sequences derived from samples containing *S. epidermidis* strains, and comparison with the present sequence allows for the identification of sequences that can be used to discriminate between strains, as well as those that are common to all *S. epidermidis* strains. In one embodiment, the invention provides nucleic acids, including probes, and peptide and polypeptide sequences that discriminate between different strains of *S. epidermidis*. Strain-specific components can also be identified functionally by their ability to elicit or react with antibodies that selectively recognize one or more *S. epidermidis* strains.

In another embodiment, the invention provides nucleic acids, including probes, and peptide and polypeptide sequences that are common to all *S. epidermidis* strains but are not found in other bacterial species.

S. epidermidis Polypeptides

This invention encompasses isolated *S. epidermidis* polypeptides encoded by the disclosed *S. epidermidis* genomic sequences, including the polypeptides of the invention contained in the Sequence Listing. Polypeptides of the invention are preferably at least about 5 amino acid residues in length. Using the DNA sequence information provided herein, the amino acid sequences of the polypeptides encompassed by the invention can be deduced using methods well-known in the art. It will be understood that the sequence of an entire nucleic acid encoding an *S. epidermidis* polypeptide can be isolated and identified based on an ORF that encodes only a fragment of the cognate protein-coding region. This can be achieved, for example, by using the isolated nucleic acid encoding the ORF, or fragments thereof, to prime

a polymerase chain reaction with genomic *S. epidermidis* DNA as template; this is followed by sequencing the amplified product.

The polypeptides of the present invention, including function-conservative variants of the disclosed ORFs, may be isolated from wild-type or mutant *S. epidermidis* cells, or from heterologous organisms or cells (including, but not limited to, bacteria, fungi, insect, plant, and mammalian cells) including *S. epidermidis* into which an *S. epidermidis* -derived protein-coding sequence has been introduced and expressed. Furthermore, the polypeptides may be part of recombinant fusion proteins.

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S. epidermidis polypeptides of the invention can be chemically synthesized using commercially automated procedures such as those referenced herein, including, without limitation, exclusive solid phase synthesis, partial solid phase methods, fragment condensation or classical solution synthesis. The polypeptides are preferably prepared by solid phase peptide synthesis as described by Merrifield, 1963, J. Am. Chem. Soc. 85:2149. The synthesis is carried out with amino acids that are protected at the alpha-amino terminus.

Trifunctional amino acids with labile side-chains are also protected with suitable groups to prevent undesired chemical reactions from occurring during the assembly of the polypeptides. The alpha-amino protecting group is selectively removed to allow subsequent reaction to take place at the amino-terminus. The conditions for the removal of the alpha-amino protecting group do not remove the side-chain protecting groups.

Methods for polypeptide purification are well-known in the art, including, without limitation, preparative disc-gel electrophoresis, isoelectric focusing, HPLC, reversed-phase HPLC, gel filtration, ion exchange and partition chromatography, and countercurrent distribution. For some purposes, it is preferable to produce the polypeptide in a recombinant system in which the *S. epidermidis* protein contains an additional sequence tag that facilitates purification, such as, but not limited to, a polyhistidine sequence. The polypeptide can then be purified from a crude lysate of the host cell by chromatography on an appropriate solid-phase matrix. Alternatively, antibodies produced against an *S. epidermidis* protein or against

peptides derived therefrom can be used as purification reagents. Other purification methods are possible.

The present invention also encompasses derivatives and homologues of *S. epidermidis* -encoded polypeptides. For some purposes, nucleic acid sequences encoding the peptides may be altered by substitutions, additions, or deletions that provide for functionally equivalent molecules, i.e., function-conservative variants. For example, one or more amino acid residues within the sequence can be substituted by another amino acid of similar properties, such as, for example, positively charged amino acids (arginine, lysine, and histidine); negatively charged amino acids (aspartate and glutamate); polar neutral amino acids; and non-polar amino acids.

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The isolated polypeptides may be modified by, for example, phosphorylation, sulfation, acylation, or other protein modifications. They may also be modified with a label capable of providing a detectable signal, either directly or indirectly, including, but not limited to, radioisotopes and fluorescent compounds.

To identify *S. epidermidis* -derived polypeptides for use in the present invention, essentially the complete genomic sequence of a virulent, methicillin-resistant isolate of *Staphylococcus mirabilis* isolate was analyzed. While, in very rare instances, a nucleic acid sequencing error may be revealed, resolving a rare sequencing error is well within the art, and such an occurrence will not prevent one skilled in the art from practicing the invention.

Also encompassed are any *S. epidermidis* polypeptide sequences that are contained within the open reading frames (ORFs), including complete protein-coding sequences, of which any of SEQ ID NO: 1 - SEQ ID NO: 3772 forms a part. Table 2, which is appended herewith and which forms part of the present specification, provides a putative identification of the particular function of a polypeptide which is encoded by each ORF, based on the homology match (determined by the BLAST algorithm) of the predicted polypeptide with known proteins encoded by ORFs in other organisms. As a result, one skilled in the art can

use the polypeptides of the present invention for commercial and industrial purposes consistent with the type of putative identification of the polypeptide.

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The present invention provides a library of *S. epidermidis* -derived polypeptide sequences, and a corresponding library of nucleic acid sequences encoding the polypeptides, wherein the polypeptides themselves, or polypeptides contained within ORFs of which they form a part, comprise sequences that are contemplated for use as components of vaccines. Non-limiting examples of such sequences are listed by SEQ ID NO in Table 2, which is appended herewith and which forms part of the present specification.

The present invention also provides a library of *S. epidermidis* -derived polypeptide sequences, and a corresponding library of nucleic acid sequences encoding the polypeptides, wherein the polypeptides themselves, or polypeptides contained within ORFs of which they form a part, comprise sequences lacking homology to any known prokaryotic or eukaryotic sequences. Such libraries provide probes, primers, and markers which can be used to diagnose *S. epidermidis* infection, including use as markers in epidemiological studies. Non-limiting examples of such sequences are listed by SEQ ID NO in Table 2, which is appended hereto and part hereof.

The present invention also provides a library of *S. epidermidis* -derived polypeptide sequences, and a corresponding library of nucleic acid sequences encoding the polypeptides, wherein the polypeptides themselves, or polypeptides contained within ORFs of which they form a part, comprise targets for therapeutic drugs.

Specific Example: Determination Of Staphylococcus Protein Antigens For Antibody And Vaccine Development

The selection of Staphylococcus protein antigens for vaccine development can be

derived from the nucleic acids encoding *S. epidermidis* polypeptides. First, the ORF's can be
analyzed for homology to other known exported or membrane proteins and analyzed using the
discriminant analysis described by Klein, et al. (Klein, P., Kanehsia, M., and DeLisi, C.

(1985) Biochimica et Biophysica Acta 815, 468-476) for predicting exported and membrane proteins.

Homology searches can be performed using the BLAST algorithm contained in the Wisconsin Sequence Analysis Package (Genetics Computer Group, University Research Park, 575 Science Drive, Madison, WI 53711) to compare each predicted ORF amino acid sequence with all sequences found in the current GenBank, SWISS-PROT and PIR databases. BLAST searches for local alignments between the ORF and the databank sequences and reports a probability score which indicates the probability of finding this sequence by chance in the database. ORF's with significant homology (e.g. probabilities lower than 1x10 that the

Discriminant analysis (Klein, et al. supra) can be used to examine the ORF amino acid sequences. This algorithm uses the intrinsic information contained in the ORF amino acid sequence and compares it to information derived from the properties of known membrane and exported proteins. This comparison predicts which proteins will be exported, membrane associated or cytoplasmic. ORF amino acid sequences identified as exported or membrane associated by this algorithm are likely protein antigens for vaccine development.

20 Production of Fragments and Analogs of S. epidermidis Nucleic Acids and Polypeptides

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Based on the discovery of the *S. epidermidis* gene products of the invention provided in the Sequence Listing, one skilled in the art can alter the disclosed structure of *S. epidermidis* genes, e.g., by producing fragments or analogs, and test the newly produced structures for activity. Examples of techniques known to those skilled in the relevant art which allow the production and testing of fragments and analogs are discussed below. These, or analogous methods can be used to make and screen libraries of polypeptides, e.g., libraries of random peptides or libraries of fragments or analogs of cellular proteins for the ability to

bind S. epidermidis polypeptides. Such screens are useful for the identification of inhibitors of S. epidermidis.

Generation of Fragments

Fragments of a protein can be produced in several ways, e.g., recombinantly, by proteolytic digestion, or by chemical synthesis. Internal or terminal fragments of a polypeptide can be generated by removing one or more nucleotides from one end (for a terminal fragment) or both ends (for an internal fragment) of a nucleic acid which encodes the polypeptide. Expression of the mutagenized DNA produces polypeptide fragments.

10 Digestion with "end-nibbling" endonucleases can thus generate DNAs which encode an array of fragments. DNAs which encode fragments of a protein can also be generated by random shearing, restriction digestion or a combination of the above-discussed methods.

Fragments can also be chemically synthesized using techniques known in the art such as conventional Merrifield solid phase f-Moc or t-Boc chemistry. For example, peptides of the present invention may be arbitrarily divided into fragments of desired length with no overlap of the fragments, or divided into overlapping fragments of a desired length.

Alteration of Nucleic Acids and Polypeptides: Random Methods

Amino acid sequence variants of a protein can be prepared by random mutagenesis of DNA which encodes a protein or a particular domain or region of a protein. Useful methods include PCR mutagenesis and saturation mutagenesis. A library of random amino acid sequence variants can also be generated by the synthesis of a set of degenerate oligonucleotide sequences. (Methods for screening proteins in a library of variants are elsewhere herein).

25 PCR Mutagenesis

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In PCR mutagenesis, reduced Taq polymerase fidelity is used to introduce random mutations into a cloned fragment of DNA (Leung et al., 1989, *Technique* 1:11-15). The DNA

region to be mutagenized is amplified using the polymerase chain reaction (PCR) under conditions that reduce the fidelity of DNA synthesis by Taq DNA polymerase, e.g., by using a dGTP/dATP ratio of five and adding Mn to the PCR reaction. The pool of amplified DNA fragments are inserted into appropriate cloning vectors to provide random mutant libraries.

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Saturation Mutagenesis

Saturation mutagenesis allows for the rapid introduction of a large number of single base substitutions into cloned DNA fragments (Mayers et al., 1985, *Science* 229:242). This technique includes generation of mutations, e.g., by chemical treatment or irradiation of single-stranded DNA *in vitro*, and synthesis of a complimentary DNA strand. The mutation frequency can be modulated by modulating the severity of the treatment, and essentially all possible base substitutions can be obtained. Because this procedure does not involve a genetic selection for mutant fragments both neutral substitutions, as well as those that alter function, are obtained. The distribution of point mutations is not biased toward conserved sequence elements.

Degenerate Oligonucleotides

A library of homologs can also be generated from a set of degenerate oligonucleotide sequences. Chemical synthesis of a degenerate sequences can be carried out in an automatic DNA synthesizer, and the synthetic genes then ligated into an appropriate expression vector. The synthesis of degenerate oligonucleotides is known in the art (see for example, Narang, SA (1983) Tetrahedron 39:3; Itakura et al. (1981) Recombinant DNA, Proc 3rd Cleveland Sympos. Macromolecules, ed. AG Walton, Amsterdam: Elsevier pp273-289; Itakura et al. (1984) Annu. Rev. Biochem. 53:323; Itakura et al. (1984) Science 198:1056; Ike et al. (1983) Nucleic Acid Res. 11:477. Such techniques have been employed in the directed evolution of other proteins (see, for example, Scott et al. (1990) Science 249:386-390; Roberts et al. (1992)

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PNAS 89:2429-2433; Devlin et al. (1990) *Science* 249: 404-406; Cwirla et al. (1990) *PNAS* 87: 6378-6382; as well as U.S. Patents Nos. 5,223,409, 5,198,346, and 5,096,815).

Alteration of Nucleic Acids and Polypeptides: Methods for Directed Mutagenesis

Non-random or directed, mutagenesis techniques can be used to provide specific sequences or mutations in specific regions. These techniques can be used to create variants which include, e.g., deletions, insertions, or substitutions, of residues of the known amino acid sequence of a protein. The sites for mutation can be modified individually or in series, e.g., by (1) substituting first with conserved amino acids and then with more radical choices depending upon results achieved, (2) deleting the target residue, or (3) inserting residues of the same or a different class adjacent to the located site, or combinations of options 1-3.

Alanine Scanning Mutagenesis

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Alanine scanning mutagenesis is a useful method for identification of certain residues or regions of the desired protein that are preferred locations or domains for mutagenesis, Cunningham and Wells (*Science* 244:1081-1085, 1989). In alanine scanning, a residue or group of target residues are identified (e.g., charged residues such as Arg, Asp, His, Lys, and Glu) and replaced by a neutral or negatively charged amino acid (most preferably alanine or polyalanine). Replacement of an amino acid can affect the interaction of the amino acids with the surrounding aqueous environment in or outside the cell. Those domains demonstrating functional sensitivity to the substitutions are then refined by introducing further or other variants at or for the sites of substitution. Thus, while the site for introducing an amino acid sequence variation is predetermined, the nature of the mutation per se need not be predetermined. For example, to optimize the performance of a mutation at a given site, alanine scanning or random mutagenesis may be conducted at the target codon or region and the expressed desired protein subunit variants are screened for the optimal combination of desired activity.

Oligonucleotide-Mediated Mutagenesis

Oligonucleotide-mediated mutagenesis is a useful method for preparing substitution, deletion, and insertion variants of DNA, see, e.g., Adelman et al., (*DNA* 2:183, 1983).

Briefly, the desired DNA is altered by hybridizing an oligonucleotide encoding a mutation to a DNA template, where the template is the single-stranded form of a plasmid or bacteriophage containing the unaltered or native DNA sequence of the desired protein. After hybridization, a DNA polymerase is used to synthesize an entire second complementary strand of the template that will thus incorporate the oligonucleotide primer, and will code for the selected alteration in the desired protein DNA. Generally, oligonucleotides of at least about 25 nucleotides in length are used. An optimal oligonucleotide will have 12 to 15 nucleotides that are completely complementary to the template on either side of the nucleotide(s) coding for the mutation. This ensures that the oligonucleotide will hybridize properly to the single-stranded DNA template molecule. The oligonucleotides are readily synthesized using techniques known in the art such as that described by Crea et al. (*Proc. Natl. Acad. Sci.* USA, 75: 5765[1978]).

Cassette Mutagenesis

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Another method for preparing variants, cassette mutagenesis, is based on the technique described by Wells et al. (*Gene*, 34:315[1985]). The starting material is a plasmid (or other vector) which includes the protein subunit DNA to be mutated. The codon(s) in the protein subunit DNA to be mutated are identified. There must be a unique restriction endonuclease site on each side of the identified mutation site(s). If no such restriction sites exist, they may be generated using the above-described oligonucleotide-mediated mutagenesis method to introduce them at appropriate locations in the desired protein subunit DNA. After the restriction sites have been introduced into the plasmid, the plasmid is cut at these sites to linearize it. A double-stranded oligonucleotide encoding the sequence of the DNA between

the restriction sites but containing the desired mutation(s) is synthesized using standard procedures. The two strands are synthesized separately and then hybridized together using standard techniques. This double-stranded oligonucleotide is referred to as the cassette. This cassette is designed to have 3' and 5' ends that are comparable with the ends of the linearized plasmid, such that it can be directly ligated to the plasmid. This plasmid now contains the mutated desired protein subunit DNA sequence.

Combinatorial Mutagenesis

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Combinatorial mutagenesis can also be used to generate mutants (Ladner et al., WO 88/06630). In this method, the amino acid sequences for a group of homologs or other related proteins are aligned, preferably to promote the highest homology possible. All of the amino acids which appear at a given position of the aligned sequences can be selected to create a degenerate set of combinatorial sequences. The variegated library of variants is generated by combinatorial mutagenesis at the nucleic acid level, and is encoded by a variegated gene library. For example, a mixture of synthetic oligonucleotides can be enzymatically ligated into gene sequences such that the degenerate set of potential sequences are expressible as individual peptides, or alternatively, as a set of larger fusion proteins containing the set of degenerate sequences.

20 Other Modifications of S. epidermidis Nucleic Acids and Polypeptides

It is possible to modify the structure of an *S. epidermidis* polypeptide for such purposes as increasing solubility, enhancing stability (e.g., shelf life *ex vivo* and resistance to proteolytic degradation *in vivo*). A modified *S. epidermidis* protein or peptide can be produced in which the amino acid sequence has been altered, such as by amino acid substitution, deletion, or addition as described herein.

An S. epidermidis peptide can also be modified by substitution of cysteine residues preferably with alanine, serine, threonine, leucine or glutamic acid residues to minimize

dimerization via disulfide linkages. In addition, amino acid side chains of fragments of the protein of the invention can be chemically modified. Another modification is cyclization of the peptide.

In order to enhance stability and/or reactivity, an S. epidermidis polypeptide can be modified to incorporate one or more polymorphisms in the amino acid sequence of the protein 5 resulting from any natural allelic variation. Additionally, D-amino acids, non-natural amino acids, or non-amino acid analogs can be substituted or added to produce a modified protein within the scope of this invention. Furthermore, an S. epidermidis polypeptide can be modified using polyethylene glycol (PEG) according to the method of A. Sehon and co-10 workers (Wie et al., supra) to produce a protein conjugated with PEG. In addition, PEG can be added during chemical synthesis of the protein. Other modifications of S. epidermidis proteins include reduction/alkylation (Tarr, Methods of Protein Microcharacterization, J. E. Silver ed., Humana Press, Clifton NJ 155-194 (1986)); acylation (Tarr, supra); chemical coupling to an appropriate carrier (Mishell and Shiigi, eds, Selected Methods in Cellular 15 Immunology, WH Freeman, San Francisco, CA (1980), U.S. Patent 4,939,239; or mild formalin treatment (Marsh, (1971) Int. Arch. of Allergy and Appl. Immunol., 41: 199 - 215).

To facilitate purification and potentially increase solubility of an *S. epidermidis* protein or peptide, it is possible to add an amino acid fusion moiety to the peptide backbone. For example, hexa-histidine can be added to the protein for purification by immobilized metal ion affinity chromatography (Hochuli, E. et al., (1988) *Bio/Technology*, 6: 1321 - 1325). In addition, to facilitate isolation of peptides free of irrelevant sequences, specific endoprotease cleavage sites can be introduced between the sequences of the fusion moiety and the peptide.

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To potentially aid proper antigen processing of epitopes within an *S. epidermidis* polypeptide, canonical protease sensitive sites can be engineered between regions, each comprising at least one epitope via recombinant or synthetic methods. For example, charged amino acid pairs, such as KK or RR, can be introduced between regions within a protein or fragment during recombinant construction thereof. The resulting peptide can be rendered

sensitive to cleavage by cathepsin and/or other trypsin-like enzymes which would generate portions of the protein containing one or more epitopes. In addition, such charged amino acid residues can result in an increase in the solubility of the peptide.

5 Primary Methods for Screening Polypeptides and Analogs

Various techniques are known in the art for screening generated mutant gene products. Techniques for screening large gene libraries often include cloning the gene library into replicable expression vectors, transforming appropriate cells with the resulting library of vectors, and expressing the genes under conditions in which detection of a desired activity, e.g., in this case, binding to *S. epidermidis* polypeptide or an interacting protein, facilitates relatively easy isolation of the vector encoding the gene whose product was detected. Each of the techniques described below is amenable to high through-put analysis for screening large numbers of sequences created, e.g., by random mutagenesis techniques.

15 <u>Two Hybrid Systems</u>

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Two hybrid assays such as the system described below (as with the other screening methods described herein), can be used to identify polypeptides, e.g., fragments or analogs of a naturally-occurring *S. epidermidis* polypeptide, e.g., of cellular proteins, or of randomly generated polypeptides which bind to an *S. epidermidis* protein. (The *S. epidermidis* domain is used as the bait protein and the library of variants are expressed as prey fusion proteins.) In an analogous fashion, a two hybrid assay (as with the other screening methods described herein), can be used to find polypeptides which bind an *S. epidermidis* polypeptide.

Display Libraries

In one approach to screening assays, the Staphylococcus peptides are displayed on the surface of a cell or viral particle, and the ability of particular cells or viral particles to bind an appropriate receptor protein via the displayed product is detected in a "panning assay". For

example, the gene library can be cloned into the gene for a surface membrane protein of a bacterial cell, and the resulting fusion protein detected by panning (Ladner et al., WO 88/06630; Fuchs et al. (1991) *Bio/Technology* 9:1370-1371; and Goward et al. (1992) *TIBS* 18:136-140). In a similar fashion, a detectably labeled ligand can be used to score for potentially functional peptide homologs. Fluorescently labeled ligands, e.g., receptors, can be used to detect homologs which retain ligand-binding activity. The use of fluorescently labeled ligands, allows cells to be visually inspected and separated under a fluorescence microscope, or, where the morphology of the cell permits, to be separated by a fluorescence-activated cell sorter.

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A gene library can be expressed as a fusion protein on the surface of a viral particle. For instance, in the filamentous phage system, foreign peptide sequences can be expressed on the surface of infectious phage, thereby conferring two significant benefits. First, since these phage can be applied to affinity matrices at concentrations well over 10 hage per milliliter, a large number of phage can be screened at one time. Second, since each infectious phage displays a gene product on its surface, if a particular phage is recovered from an affinity matrix in low yield, the phage can be amplified by another round of infection. The group of almost identical *E. coli* filamentous phages, M13, fd., and f1, are most often used in phage display libraries. Either of the phage gIII or gVIII coat proteins can be used to generate fusion proteins without disrupting the ultimate packaging of the viral particle. Foreign epitopes can be expressed at the NH2-terminal end of pIII and phage bearing such epitopes recovered from a large excess of phage lacking this epitope (Ladner et al. PCT publication WO 90/02909; Garrard et al., PCT publication WO 92/09690; Marks et al. (1992) *J. Biol. Chem.* 267:16007-16010; Griffiths et al. (1993) *EMBO J* 12:725-734; Clackson et al. (1991) *Nature* 352:624-628; and Barbas et al. (1992) *PNAS* 89:4457-4461).

A common approach uses the maltose receptor of *E. coli* (the outer membrane protein, LamB) as a peptide fusion partner (Charbit et al. (1986) *EMBO* 5, 3029-3037).

Oligonucleotides have been inserted into plasmids encoding the LamB gene to produce

peptides fused into one of the extracellular loops of the protein. These peptides are available for binding to ligands, e.g., to antibodies, and can elicit an immune response when the cells are administered to animals. Other cell surface proteins, e.g., OmpA (Schorr et al. (1991) Vaccines 91, pp. 387-392), PhoE (Agterberg, et al. (1990) Gene 88, 37-45), and PAL (Fuchs et al. (1991) Bio/Tech 9, 1369-1372), as well as large bacterial surface structures have served as vehicles for peptide display. Peptides can be fused to pilin, a protein which polymerizes to form the pilus-a conduit for interbacterial exchange of genetic information (Thiry et al. (1989) Appl. Environ. Microbiol. 55, 984-993). Because of its role in interacting with other cells, the pilus provides a useful support for the presentation of peptides to the extracellular environment. Another large surface structure used for peptide display is the bacterial motive organ, the flagellum. Fusion of peptides to the subunit protein flagellin offers a dense array of many peptide copies on the host cells (Kuwajima et al. (1988) *Bio/Tech.* 6, 1080-1083). Surface proteins of other bacterial species have also served as peptide fusion partners. Examples include the Staphylococcus protein A and the outer membrane IgA protease of Neisseria (Hansson et al. (1992) J. Bacteriol. 174, 4239-4245 and Klauser et al. (1990) EMBO J. 9, 1991-1999).

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In the filamentous phage systems and the LamB system described above, the physical link between the peptide and its encoding DNA occurs by the containment of the DNA within a particle (cell or phage) that carries the peptide on its surface. Capturing the peptide captures the particle and the DNA within. An alternative scheme uses the DNA-binding protein LacI to form a link between peptide and DNA (Cull *et al.* (1992) *PNAS USA* 89:1865-1869). This system uses a plasmid containing the LacI gene with an oligonucleotide cloning site at its 3'-end. Under the controlled induction by arabinose, a LacI-peptide fusion protein is produced. This fusion retains the natural ability of LacI to bind to a short DNA sequence known as LacO operator (LacO). By installing two copies of LacO on the expression plasmid, the LacI-peptide fusion binds tightly to the plasmid that encoded it. Because the plasmids in each cell contain only a single oligonucleotide sequence and each cell expresses only a single peptide

sequence, the peptides become specifically and stablely associated with the DNA sequence that directed its synthesis. The cells of the library are gently lysed and the peptide-DNA complexes are exposed to a matrix of immobilized receptor to recover the complexes containing active peptides. The associated plasmid DNA is then reintroduced into cells for amplification and DNA sequencing to determine the identity of the peptide ligands. As a demonstration of the practical utility of the method, a large random library of dodecapeptides was made and selected on a monoclonal antibody raised against the opioid peptide dynorphin B. A cohort of peptides was recovered, all related by a consensus sequence corresponding to a six-residue portion of dynorphin B. (Cull et al. (1992) *Proc. Natl. Acad. Sci. U.S.*A. 89-1869)

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This scheme, sometimes referred to as peptides-on-plasmids, differs in two important ways from the phage display methods. First, the peptides are attached to the C-terminus of the fusion protein, resulting in the display of the library members as peptides having free carboxy termini. Both of the filamentous phage coat proteins, pIII and pVIII, are anchored to the phage through their C-termini, and the guest peptides are placed into the outward-extending N-terminal domains. In some designs, the phage-displayed peptides are presented right at the amino terminus of the fusion protein. (Cwirla, et al. (1990) Proc. Natl. Acad. Sci. U.S.A. 87, 6378-6382) A second difference is the set of biological biases affecting the population of peptides actually present in the libraries. The LacI fusion molecules are confined to the cytoplasm of the host cells. The phage coat fusions are exposed briefly to the cytoplasm during translation but are rapidly secreted through the inner membrane into the periplasmic compartment, remaining anchored in the membrane by their C-terminal hydrophobic domains. with the N-termini, containing the peptides, protruding into the periplasm while awaiting assembly into phage particles. The peptides in the LacI and phage libraries may differ significantly as a result of their exposure to different proteolytic activities. The phage coat proteins require transport across the inner membrane and signal peptidase processing as a prelude to incorporation into phage. Certain peptides exert a deleterious effect on these

processes and are underrepresented in the libraries (Gallop et al. (1994) *J. Med. Chem.* 37(9):1233-1251). These particular biases are not a factor in the LacI display system.

The number of small peptides available in recombinant random libraries is enormous.

7 19 independent clones are routinely prepared. Libraries as large as 10 11 recombinants have been created, but this size approaches the practical limit for clone libraries. This limitation in library size occurs at the step of transforming the DNA containing randomized segments into the host bacterial cells. To circumvent this limitation, an *in vitro* system based on the display of nascent peptides in polysome complexes has recently been developed. This display library method has the potential of producing libraries 3-6 orders of magnitude larger than the currently available phage/phagemid or plasmid libraries.

Furthermore, the construction of the libraries, expression of the peptides, and screening, is done in an entirely cell-free format.

In one application of this method (Gallop et al. (1994) *J. Med. Chem.* 37(9):1233-1251), a molecular DNA library encoding 10 decapeptides was constructed and the library expressed in an *E. coli* S30 *in vitro* coupled transcription/translation system. Conditions were chosen to stall the ribosomes on the mRNA, causing the accumulation of a substantial proportion of the RNA in polysomes and yielding complexes containing nascent peptides still linked to their encoding RNA. The polysomes are sufficiently robust to be affinity purified on immobilized receptors in much the same way as the more conventional recombinant peptide display libraries are screened. RNA from the bound complexes is recovered, converted to cDNA, and amplified by PCR to produce a template for the next round of synthesis and screening. The polysome display method can be coupled to the phage display system. Following several rounds of screening, cDNA from the enriched pool of polysomes was cloned into a phagemid vector. This vector serves as both a peptide expression vector, displaying peptides fused to the coat proteins, and as a DNA sequencing vector for peptide identification. By expressing the polysome-derived peptides on phage, one can either continue the affinity selection procedure in this format or assay the peptides on individual

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clones for binding activity in a phage ELISA, or for binding specificity in a completion phage ELISA (Barret, et al. (1992) *Anal. Biochem* 204,357-364). To identify the sequences of the active peptides one sequences the DNA produced by the phagemid host.

5 Secondary Screening of Polypeptides and Analogs

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The high through-put assays described above can be followed by secondary screens in order to identify further biological activities which will, e.g., allow one skilled in the art to differentiate agonists from antagonists. The type of a secondary screen used will depend on the desired activity that needs to be tested. For example, an assay can be developed in which the ability to inhibit an interaction between a protein of interest and its respective ligand can be used to identify antagonists from a group of peptide fragments isolated though one of the primary screens described above.

Therefore, methods for generating fragments and analogs and testing them for activity are known in the art. Once the core sequence of interest is identified, it is routine for one skilled in the art to obtain analogs and fragments.

Peptide Mimetics of S. epidermidis Polypeptides

The invention also provides for reduction of the protein binding domains of the subject *S. epidermidis* polypeptides to generate mimetics, e.g. peptide or non-peptide agents. The peptide mimetics are able to disrupt binding of a polypeptide to its counter ligand, e.g., in the case of an *S. epidermidis* polypeptide binding to a naturally occurring ligand. The critical residues of a subject *S. epidermidis* polypeptide which are involved in molecular recognition of a polypeptide can be determined and used to generate *S. epidermidis* -derived peptidomimetics which competitively or noncompetitively inhibit binding of the *S. epidermidis* polypeptide with an interacting polypeptide (see, for example, European patent applications EP-412,762A and EP-B31,080A).

For example, scanning mutagenesis can be used to map the amino acid residues of a particular S. epidermidis polypeptide involved in binding an interacting polypeptide, peptidomimetic compounds (e.g. diazepine or isoquinoline derivatives) can be generated which mimic those residues in binding to an interacting polypeptide, and which therefore can inhibit binding of an S. epidermidis polypeptide to an interacting polypeptide and thereby 5 interfere with the function of S. epidermidis polypeptide. For instance, non-hydrolyzable peptide analogs of such residues can be generated using benzodiazepine (e.g., see Freidinger et al. in *Peptides: Chemistry and Biology*, G.R. Marshall ed., ESCOM Publisher: Leiden, Netherlands, 1988), azepine (e.g., see Huffman et al. in *Peptides: Chemistry and Biology*, 10 G.R. Marshall ed., ESCOM Publisher: Leiden, Netherlands, 1988), substituted gama lactam rings (Garvey et al. in *Peptides: Chemistry and Biology*, G.R. Marshall ed., ESCOM Publisher: Leiden, Netherlands, 1988), keto-methylene pseudopeptides (Ewenson et al. (1986) J Med Chem 29:295; and Ewenson et al. in Peptides: Structure and Function (Proceedings of the 9th American Peptide Symposium) Pierce Chemical Co. Rockland, IL, 1985), b-turn 15 dipeptide cores (Nagai et al. (1985) Tetrahedron Lett 26:647; and Sato et al. (1986) J Chem Soc Perkin Trans 1:1231), and b-aminoalcohols (Gordon et al. (1985) Biochem Biophys Res Commun 126:419; and et al. (1986) Biochem Biophys Res Commun 134:71).

Vaccine Formulations for S. epidermidis Nucleic Acids and Polypeptides

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This invention also features vaccine compositions for protection against infection by *S. epidermidis* or for treatment of *S. epidermidis* infection. In one embodiment, the vaccine compositions contain one or more immunogenic components such as a surface protein from *S. epidermidis*, or portion thereof, and a pharmaceutically acceptable carrier. Nucleic acids within the scope of the invention are exemplified by the nucleic acids of the invention contained in the Sequence Listing which encode *S. epidermidis* surface proteins. Any nucleic acid encoding an immunogenic *S. epidermidis* protein, or portion thereof, which is capable of

expression in a cell, can be used in the present invention. These vaccines have therapeutic and prophylactic utilities.

One aspect of the invention provides a vaccine composition for protection against infection by *S. epidermidis* which contains at least one immunogenic fragment of an *S. epidermidis* protein and a pharmaceutically acceptable carrier. Preferred fragments include peptides of at least about 10 amino acid residues in length, preferably about 10-20 amino acid residues in length, and more preferably about 12-16 amino acid residues in length.

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Immunogenic components of the invention can be obtained, for example, by screening polypeptides recombinantly produced from the corresponding fragment of the nucleic acid encoding the full-length *S. epidermidis* protein. In addition, fragments can be chemically synthesized using techniques known in the art such as conventional Merrifield solid phase f-Moc or t-Boc chemistry.

In one embodiment, immunogenic components are identified by the ability of the peptide to stimulate T cells. Peptides which stimulate T cells, as determined by, for example, T cell proliferation or cytokine secretion are defined herein as comprising at least one T cell epitope. T cell epitopes are believed to be involved in initiation and perpetuation of the immune response to the protein allergen which is responsible for the clinical symptoms of allergy. These T cell epitopes are thought to trigger early events at the level of the T helper cell by binding to an appropriate HLA molecule on the surface of an antigen presenting cell, thereby stimulating the T cell subpopulation with the relevant T cell receptor for the epitope. These events lead to T cell proliferation, lymphokine secretion, local inflammatory reactions, recruitment of additional immune cells to the site of antigen/T cell interaction, and activation of the B cell cascade, leading to the production of antibodies. A T cell epitope is the basic element, or smallest unit of recognition by a T cell receptor, where the epitope comprises amino acids essential to receptor recognition (e.g., approximately 6 or 7 amino acid residues). Amino acid sequences which mimic those of the T cell epitopes are within the scope of this invention.

Screening immunogenic components can be accomplished using one or more of several different assays. For example, *in vitro*, peptide T cell stimulatory activity is assayed by contacting a peptide known or suspected of being immunogenic with an antigen presenting cell which presents appropriate MHC molecules in a T cell culture. Presentation of an immunogenic *S. epidermidis* peptide in association with appropriate MHC molecules to T cells in conjunction with the necessary co-stimulation has the effect of transmitting a signal to the T cell that induces the production of increased levels of cytokines, particularly of interleukin-2 and interleukin-4. The culture supernatant can be obtained and assayed for interleukin-2 or other known cytokines. For example, any one of several conventional assays for interleukin-2 can be employed, such as the assay described in *Proc. Natl. Acad. Sci USA*, 86: 1333 (1989) the pertinent portions of which are incorporated herein by reference. A kit for an assay for the production of interferon is also available from Genzyme Corporation (Cambridge, MA).

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Alternatively, a common assay for T cell proliferation entails measuring tritiated thymidine incorporation. The proliferation of T cells can be measured *in vitro* by determining the amount of ³H-labeled thymidine incorporated into the replicating DNA of cultured cells. Therefore, the rate of DNA synthesis and, in turn, the rate of cell division can be quantified.

Vaccine compositions of the invention containing immunogenic components (e.g., *S. epidermidis* polypeptide or fragment thereof or nucleic acid encoding an *S. epidermidis* polypeptide or fragment thereof) preferably include a pharmaceutically acceptable carrier. The term "pharmaceutically acceptable carrier" refers to a carrier that does not cause an allergic reaction or other untoward effect in patients to whom it is administered. Suitable pharmaceutically acceptable carriers include, for example, one or more of water, saline, phosphate buffered saline, dextrose, glycerol, ethanol and the like, as well as combinations thereof. Pharmaceutically acceptable carriers may further comprise minor amounts of auxiliary substances such as wetting or emulsifying agents, preservatives or buffers, which enhance the shelf life or effectiveness of the antibody. For vaccines of the invention

containing *S. epidermidis* polypeptides, the polypeptide is co-administered with a suitable adjuvant.

It will be apparent to those of skill in the art that the therapeutically effective amount of DNA or protein of this invention will depend, *inter alia*, upon the administration schedule, the unit dose of antibody administered, whether the protein or DNA is administered in combination with other therapeutic agents, the immune status and health of the patient, and the therapeutic activity of the particular protein or DNA.

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Vaccine compositions are conventionally administered parenterally, e.g., by injection, either subcutaneously or intramuscularly. Methods for intramuscular immunization are described by Wolff et al. (1990) *Science* 247: 1465-1468 and by Sedegah et al. (1994) *Immunology* 91: 9866-9870. Other modes of administration include oral and pulmonary formulations, suppositories, and transdermal applications. Oral immunization is preferred over parenteral methods for inducing protection against infection by *S. epidermidis*. Cain et. al. (1993) *Vaccine* 11: 637-642. Oral formulations include such normally employed excipients as, for example, pharmaceutical grades of mannitol, lactose, starch, magnesium stearate, sodium saccharine, cellulose, magnesium carbonate, and the like.

The vaccine compositions of the invention can include an adjuvant, including, but not limited to aluminum hydroxide; N-acetyl-muramyl--L-threonyl-D-isoglutamine (thr-MDP); N-acetyl-nor-muramyl-L-alanyl-D-isoglutamine (CGP 11637, referred to as nor-MDP); N-acetylmuramyl-L-alanyl-D-isoglutaminyl-L-alanine-2-(1'-2'-dipalmitoyl-sn-glycero-3-hydroxyphos-phoryloxy)-ethylamine (CGP 19835A, referred to a MTP-PE); RIBI, which contains three components from bacteria; monophosphoryl lipid A; trehalose dimycoloate; cell wall skeleton (MPL + TDM + CWS) in a 2% squalene/Tween 80 emulsion; and cholera toxin. Others which may be used are non-toxic derivatives of cholera toxin, including its B subunit, and/or conjugates or genetically engineered fusions of the *S. epidermidis* polypeptide with cholera toxin or its B subunit, procholeragenoid, fungal polysaccharides, including

schizophyllan, muramyl dipeptide, muramyl dipeptide derivatives, phorbol esters, labile toxin of *E. coli*, non-*S. epidermidis* bacterial lysates, block polymers or saponins.

Other suitable delivery methods include biodegradable microcapsules or immunostimulating complexes (ISCOMs), cochleates, or liposomes, genetically engineered attenuated live vectors such as viruses or bacteria, and recombinant (chimeric) virus-like particles, e.g., bluetongue. The amount of adjuvant employed will depend on the type of adjuvant used. For example, when the mucosal adjuvant is cholera toxin, it is suitably used in an amount of 5 mg to 50 mg, for example 10 mg to 35 mg. When used in the form of microcapsules, the amount used will depend on the amount employed in the matrix of the microcapsule to achieve the desired dosage. The determination of this amount is within the skill of a person of ordinary skill in the art.

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Carrier systems in humans may include enteric release capsules protecting the antigen from the acidic environment of the stomach, and including *S. epidermidis* polypeptide in an insoluble form as fusion proteins. Suitable carriers for the vaccines of the invention are enteric coated capsules and polylactide-glycolide microspheres. Suitable diluents are 0.2 N NaHCO₃ and/or saline.

Vaccines of the invention can be administered as a primary prophylactic agent in adults or in children, as a secondary prevention, after successful eradication of *S. epidermidis* in an infected host, or as a therapeutic agent in the aim to induce an immune response in a susceptible host to prevent infection by *S. epidermidis*. The vaccines of the invention are administered in amounts readily determined by persons of ordinary skill in the art. Thus, for adults a suitable dosage will be in the range of 10 mg to 10 g, preferably 10 mg to 100 mg. A suitable dosage for adults will also be in the range of 5 mg to 500 mg. Similar dosage ranges will be applicable for children. Those skilled in the art will recognize that the optimal dose may be more or less depending upon the patient's body weight, disease, the route of administration, and other factors. Those skilled in the art will also recognize that appropriate dosage levels can be obtained based on results with known oral vaccines such as, for example,

a vaccine based on an *E. coli* lysate (6 mg dose daily up to total of 540 mg) and with an enterotoxigenic *E. coli* purified antigen (4 doses of 1 mg) (Schulman et al., *J. Urol.* 150:917-921 (1993); Boedecker et al., *American Gastroenterological Assoc.* 999:A-222 (1993)). The number of doses will depend upon the disease, the formulation, and efficacy data from clinical trials. Without intending any limitation as to the course of treatment, the treatment can be administered over 3 to 8 doses for a primary immunization schedule over 1 month (Boedeker, *American Gastroenterological Assoc.* 888:A-222 (1993)).

In a preferred embodiment, a vaccine composition of the invention can be based on a killed whole *E. coli* preparation with an immunogenic fragment of an *S. epidermidis* protein of the invention expressed on its surface or it can be based on an *E. coli* lysate, wherein the killed *E. coli* acts as a carrier or an adjuvant.

It will be apparent to those skilled in the art that some of the vaccine compositions of the invention are useful only for preventing *S. epidermidis* infection, some are useful only for treating *S. epidermidis* infection, and some are useful for both preventing and treating *S. epidermidis* infection. In a preferred embodiment, the vaccine composition of the invention provides protection against *S. epidermidis* infection by stimulating humoral and/or cell-mediated immunity against *S. epidermidis*. It should be understood that amelioration of any of the symptoms of *S. epidermidis* infection is a desirable clinical goal, including a lessening of the dosage of medication used to treat *S. epidermidis* -caused disease, or an increase in the production of antibodies in the serum or mucous of patients.

Antibodies Reactive With S. epidermidis Polypeptides

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The invention also includes antibodies specifically reactive with the subject *S. epidermidis* polypeptide. Anti-protein/anti-peptide antisera or monoclonal antibodies can be made by standard protocols (See, for example, *Antibodies:* A *Laboratory Manual* ed. by Harlow and Lane (Cold Spring Harbor Press: 1988)). A mammal such as a mouse, a hamster or rabbit can be immunized with an immunogenic form of the peptide. Techniques for

conferring immunogenicity on a protein or peptide include conjugation to carriers or other techniques well known in the art. An immunogenic portion of the subject *S. epidermidis* polypeptide can be administered in the presence of adjuvant. The progress of immunization can be monitored by detection of antibody titers in plasma or serum. Standard ELISA or other immunoassays can be used with the immunogen as antigen to assess the levels of antibodies.

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In a preferred embodiment, the subject antibodies are immunospecific for antigenic determinants of the *S. epidermidis* polypeptides of the invention, e.g. antigenic determinants of a polypeptide of the invention contained in the Sequence Listing, or a closely related human or non-human mammalian homolog (e.g., 90% homologous, more preferably at least about 95% homologous). In yet a further preferred embodiment of the invention, the anti-*S. epidermidis* antibodies do not substantially cross react (i.e., react specifically) with a protein which is for example, less than 80% percent homologous to a sequence of the invention contained in the Sequence Listing. By "not substantially cross react", it is meant that the antibody has a binding affinity for a non-homologous protein which is less than 10 percent, more preferably less than 5 percent, and even more preferably less than 1 percent, of the binding affinity for a protein of the invention contained in the Sequence Listing. In a most preferred embodiment, there is no cross-reactivity between bacterial and mammalian antigens.

The term antibody as used herein is intended to include fragments thereof which are also specifically reactive with *S. epidermidis* polypeptides. Antibodies can be fragmented using conventional techniques and the fragments screened for utility in the same manner as described above for whole antibodies. For example, F(ab')₂ fragments can be generated by treating antibody with pepsin. The resulting F(ab')₂ fragment can be treated to reduce disulfide bridges to produce Fab' fragments. The antibody of the invention is further intended to include bispecific and chimeric molecules having an anti-*S. epidermidis* portion.

Both monoclonal and polyclonal antibodies (Ab) directed against *S. epidermidis* polypeptides or *S. epidermidis* polypeptide variants, and antibody fragments such as Fab' and F(ab')₂, can be used to block the action of *S. epidermidis* polypeptide and allow the study of

the role of a particular *S. epidermidis* polypeptide of the invention in aberrant or unwanted intracellular signaling, as well as the normal cellular function of the *S. epidermidis* and by microinjection of anti-*S. epidermidis* polypeptide antibodies of the present invention.

Antibodies which specifically bind *S. epidermidis* epitopes can also be used in immunohistochemical staining of tissue samples in order to evaluate the abundance and pattern of expression of *S. epidermidis* antigens. Anti-*S. epidermidis* polypeptide antibodies can be used diagnostically in immuno-precipitation and immuno-blotting to detect and evaluate *S. epidermidis* levels in tissue or bodily fluid as part of a clinical testing procedure. Likewise, the ability to monitor *S. epidermidis* polypeptide levels in an individual can allow determination of the efficacy of a given treatment regimen for an individual afflicted with such a disorder. The level of an *S. epidermidis* polypeptide can be measured in cells found in bodily fluid, such as in urine samples or can be measured in tissue, such as produced by gastric biopsy. Diagnostic assays using anti-*S. epidermidis* antibodies can include, for example, immunoassays designed to aid in early diagnosis of *S. epidermidis* infections. The present invention can also be used as a method of detecting antibodies contained in samples from individuals infected by this bacterium using specific *S. epidermidis* antigens.

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Another application of anti-*S. epidermidis* polypeptide antibodies of the invention is in the immunological screening of cDNA libraries constructed in expression vectors such as λgt11, λgt18-23, λZAP, and λORF8. Messenger libraries of this type, having coding sequences inserted in the correct reading frame and orientation, can produce fusion proteins. For instance, λgt11 will produce fusion proteins whose amino termini consist of β-galactosidase amino acid sequences and whose carboxy termini consist of a foreign polypeptide. Antigenic epitopes of a subject *S. epidermidis* polypeptide can then be detected with antibodies, as, for example, reacting nitrocellulose filters lifted from infected plates with anti-*S. epidermidis* polypeptide antibodies. Phage, scored by this assay, can then be isolated from the infected plate. Thus, the presence of *S. epidermidis* gene homologs can be detected

and cloned from other species, and alternate isoforms (including splicing variants) can be detected and cloned.

Kits Containing Nucleic Acids, Polypeptides or Antibodies of the Invention

The nucleic acid, polypeptides and antibodies of the invention can be combined with other reagents and articles to form kits. Kits for diagnostic purposes typically comprise the nucleic acid, polypeptides or antibodies in vials or other suitable vessels. Kits typically comprise other reagents for performing hybridization reactions, polymerase chain reactions (PCR), or for reconstitution of lyophilized components, such as aqueous media, salts, buffers, and the like. Kits may also comprise reagents for sample processing such as detergents, chaotropic salts and the like. Kits may also comprise immobilization means such as particles, supports, wells, dipsticks and the like. Kits may also comprise labeling means such as dyes, developing reagents, radioisotopes, fluorescent agents, luminescent or chemiluminescent agents, enzymes, intercalating agents and the like. With the nucleic acid and amino acid sequence information provided herein, individuals skilled in art can readily assemble kits to serve their particular purpose. Kits further can include instructions for use.

Bio chip Technology

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The nucleic acid sequence of the present invention may be used to detect *S*.

epidermidis or other species of Staphylococcus acid sequence using bio chip technology. Bio chips containing arrays of nucleic acid sequence can also be used to measure expression of genes of *S. epidermidis* or other species of Staphylococcus. For example, to diagnose a patient with a *S. epidermidis* or other Staphylococcus infection, a sample from a human or animal can be used as a probe on a bio chip containing an array of nucleic acid sequence from the present invention. In addition, a sample from a disease state can be compared to a sample from a non-disease state which would help identify a gene that is up-regulated or expressed in the disease state. This would provide valuable insight as to the mechanism by which the

disease manifests. Changes in gene expression can also be used to identify critical pathways involved in drug transport or metabolism, and may enable the identification of novel targets involved in virulence or host cell interactions involved in maintenance of an infection. Procedures using such techniques have been described by Brown *et al.*, 1995, *Science* 270: 467-470.

Bio chips can also be used to monitor the genetic changes of potential therapeutic compounds including, deletions, insertions or mismatches. Once the therapeutic is added to the patient, changes to the genetic sequence can be evaluated for its efficacy. In addition, the nucleic acid sequence of the present invention can be used to determine essential genes in cell cycling. As described in Iyer *et al.*, 1999 (*Science*, 283:83-87) genes essential in the cell cycle can be identified using bio chips. Furthermore, the present invention provides nucleic acid sequence which can be used with bio chip technology to understand regulatory networks in bacteria, measure the response to environmental signals or drugs as in drug screening, and study virulence induction. (Mons *et al.*, 1998, *Nature Biotechnology*, 16: 45-48. Patents teaching this technology include U.S. Patents 5445934, 5744305, and 5800992.

Drug Screening Assays Using S. epidermidis Polypeptides

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By making available purified and recombinant *S. epidermidis* polypeptides, the present invention provides assays which can be used to screen for drugs which are either agonists or antagonists of the normal cellular function, in this case, of the subject *S. epidermidis* polypeptides, or of their role in intracellular signaling. Such inhibitors or potentiators may be useful as new therapeutic agents to combat *S. epidermidis* infections in humans. A variety of assay formats will suffice and, in light of the present inventions, will be comprehended by the person skilled in the art.

In many drug screening programs which test libraries of compounds and natural extracts, high throughput assays are desirable in order to maximize the number of compounds surveyed in a given period of time. Assays which are performed in cell-free systems, such as

may be derived with purified or semi-purified proteins, are often preferred as "primary" screens in that they can be generated to permit rapid development and relatively easy detection of an alteration in a molecular target which is mediated by a test compound. Moreover, the effects of cellular toxicity and/or bioavailability of the test compound can be generally ignored in the *in vitro* system, the assay instead being focused primarily on the effect of the drug on the molecular target as may be manifest in an alteration of binding affinity with other proteins or change in enzymatic properties of the molecular target. Accordingly, in an exemplary screening assay of the present invention, the compound of interest is contacted with an isolated and purified *S. epidermidis* polypeptide.

Screening assays can be constructed *in vitro* with a purified *S. epidermidis* polypeptide or fragment thereof, such as an *S. epidermidis* polypeptide having enzymatic activity, such that the activity of the polypeptide produces a detectable reaction product. The efficacy of the compound can be assessed by generating dose response curves from data obtained using various concentrations of the test compound. Moreover, a control assay can also be performed to provide a baseline for comparison. Suitable products include those with distinctive absorption, fluorescence, or chemi-luminescence properties, for example, because detection may be easily automated. A variety of synthetic or naturally occurring compounds can be tested in the assay to identify those which inhibit or potentiate the activity of the *S. epidermidis* polypeptide. Some of these active compounds may directly, or with chemical alterations to promote membrane permeability or solubility, also inhibit or potentiate the same activity (e.g., enzymatic activity) in whole, live *S. epidermidis* cells.

Overexpression Assays

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Overexpression assays are based on the premise that overproduction of a protein would lead to a higher level of resistance to compounds that selectively interfere with the function of that protein. Overexpression assays may be used to identify compounds that interfere with the function of virtually any type of protein, including without limitation

enzymes, receptors, DNA- or RNA-binding proteins, or any proteins that are directly or indirectly involved in regulating cell growth.

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Typically, two bacterial strains are constructed. One contains a single copy of the gene of interest, and a second contains several copies of the same gene. Identification of useful inhibitory compounds of this type of assay is based on a comparison of the activity of a test compound in inhibiting growth and/or viability of the two strains. The method involves constructing a nucleic acid vector that directs high level expression of a particular target nucleic acid. The vectors are then transformed into host cells in single or multiple copies to produce strains that express low to moderate and high levels of protein encoding by the target sequence (strain A and B, respectively). Nucleic acid comprising sequences encoding the target gene can, of course, be directly integrated into the host cell.

Large numbers of compounds (or crude substances which may contain active compounds) are screened for their effect on the growth of the two strains. Agents which interfere with an unrelated target equally inhibit the growth of both strains. Agents which interfere with the function of the target at high concentration should inhibit the growth of both strains. It should be possible, however, to titrate out the inhibitory effect of the compound in the overexpressing strain. That is, if the compound is affecting the particular target that is being tested, it should be possible to inhibit the growth of strain A at a concentration of the compound that allows strain B to grow.

Alternatively, a bacterial strain is constructed that contains the gene of interest under the control of an inducible promoter. Identification of useful inhibitory agents using this type of assay is based on a comparison of the activity of a test compound in inhibiting growth and/or viability of this strain under both inducing and non-inducing conditions. The method involves constructing a nucleic acid vector that directs high-level expression of a particular target nucleic acid. The vector is then transformed into host cells that are grown under both non-inducing and inducing conditions (conditions A and B, respectively).

Large numbers of compounds (or crude substances which may contain active compounds) are screened for their effect on growth under these two conditions. Agents that interfere with the function of the target should inhibit growth under both conditions. It should be possible, however, to titrate out the inhibitory effect of the compound in the overexpressing strain. That is, if the compound is affecting the particular target that is being tested, it should be possible to inhibit growth under condition A at a concentration that allows the strain to grow under condition B.

Ligand-binding Assays

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Many of the targets according to the invention have functions that have not yet been identified. Ligand-binding assays are useful to identify inhibitor compounds that interfere with the function of a particular target, even when that function is unknown. These assays are designed to detect binding of test compounds to particular targets. The detection may involve direct measurement of binding. Alternatively, indirect indications of binding may involve stabilization of protein structure or disruption of a biological function. Non-limiting examples of useful ligand-binding assays are detailed below.

A useful method for the detection and isolation of binding proteins is the Biomolecular Interaction Assay (BIAcore) system developed by Pharmacia Biosensor and described in the manufacturer's protocol (LKB Pharmacia, Sweden). The BIAcore system uses an affinity purified anti-GST antibody to immobilize GST-fusion proteins onto a sensor chip. The sensor utilizes surface plasmon resonance which is an optical phenomenon that detects changes in refractive indices. In accordance with the practice of the invention, a protein of interest is coated onto a chip and test compounds are passed over the chip. Binding is detected by a change in the refractive index (surface plasmon resonance).

A different type of ligand-binding assay involves scintillation proximity assays (SPA, described in U.S. Patent No. 4,568,649).

Another type of ligand binding assay, also undergoing development, is based on the fact that proteins containing mitochondrial targeting signals are imported into isolated mitochondria *in vitro* (Hurt *et al.*, 1985, *Embo J.* 4:2061-2068; Eilers and Schatz, *Nature*, 1986, 322:228-231). In a mitochondrial import assay, expression vectors are constructed in which nucleic acids encoding particular target proteins are inserted downstream of sequences encoding mitochondrial import signals. The chimeric proteins are synthesized and tested for their ability to be imported into isolated mitochondria in the absence and presence of test compounds. A test compound that binds to the target protein should inhibit its uptake into isolated mitochondria *in vitro*.

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Another ligand-binding assay is the yeast two-hybrid system (Fields and Song, 1989, Nature 340:245-246). The yeast two-hybrid system takes advantage of the properties of the GAL4 protein of the yeast Saccharomyces cerevisiae. The GAL4 protein is a transcriptional activator required for the expression of genes encoding enzymes of galactose utilization. This protein consists of two separable and functionally essential domains: an N-terminal domain which binds to specific DNA sequences (UAS_G); and a C-terminal domain containing acidic regions, which is necessary to activate transcription. The native GAL4 protein, containing both domains, is a potent activator of transcription when yeast are grown on galactose media. The N-terminal domain binds to DNA in a sequence-specific manner but is unable to activate transcription. The C-terminal domain contains the activating regions but cannot activate transcription because it fails to be localized to UAS_G. In the two-hybrid system, a system of two hybrid proteins containing parts of GAL4: (1) a GAL4 DNA-binding domain fused to a protein 'X' and (2) a GAL4 activation region fused to a protein 'Y'. If X and Y can form a protein-protein complex and reconstitute proximity of the GAL4 domains, transcription of a gene regulated by UAS_G occurs. Creation of two hybrid proteins, each containing one of the interacting proteins X and Y, allows the activation region of UASG to be brought to its normal site of action.

The binding assay described in Fodor *et al.*, 1991, *Science* 251:767-773, which involves testing the binding affinity of test compounds for a plurality of defined polymers synthesized on a solid substrate, may also be useful.

Compounds which bind to the polypeptides of the invention are potentially useful as antibacterial agents for use in therapeutic compositions.

Pharmaceutical formulations suitable for antibacterial therapy comprise the antibacterial agent in conjunction with one or more biologically acceptable carriers. Suitable biologically acceptable carriers include, but are not limited to, phosphate-buffered saline, saline, deionized water, or the like. Preferred biologically acceptable carriers are physiologically or pharmaceutically acceptable carriers.

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The antibacterial compositions include an antibacterial effective amount of active agent. Antibacterial effective amounts are those quantities of the antibacterial agents of the present invention that afford prophylactic protection against bacterial infections or which result in amelioration or cure of an existing bacterial infection. This antibacterial effective amount will depend upon the agent, the location and nature of the infection, and the particular host. The amount can be determined by experimentation known in the art, such as by establishing a matrix of dosages and frequencies and comparing a group of experimental units or subjects to each point in the matrix.

The antibacterial active agents or compositions can be formed into dosage unit forms, such as for example, creams, ointments, lotions, powders, liquids, tablets, capsules, suppositories, sprays, aerosols or the like. If the antibacterial composition is formulated into a dosage unit form, the dosage unit form may contain an antibacterial effective amount of active agent. Alternatively, the dosage unit form may include less than such an amount if multiple dosage unit forms or multiple dosages are to be used to administer a total dosage of the active agent. Dosage unit forms can include, in addition, one or more excipient(s), diluent(s), disintegrant(s), lubricant(s), plasticizer(s), colorant(s), dosage vehicle(s), absorption enhancer(s), stabilizer(s), bactericide(s), or the like.

For general information concerning formulations, see, e.g., Gilman et al. (eds.), 1990, Goodman and Gilman's: The Pharmacological Basis of Therapeutics, 8th ed., Pergamon Press; and Remington's Pharmaceutical Sciences, 17th ed., 1990, Mack Publishing Co., Easton, PA; Avis et al. (eds.), 1993, Pharmaceutical Dosage Forms: Parenteral Medications, Dekker, New York; Lieberman et al (eds.), 1990, Pharmaceutical Dosage Forms: Disperse Systems, Dekker, New York.

The antibacterial agents and compositions of the present invention are useful for preventing or treating *S. epidermidis* infections. Infection prevention methods incorporate a prophylactically effective amount of an antibacterial agent or composition. A prophylactically effective amount is an amount effective to prevent *S. epidermidis* infection and will depend upon the specific bacterial strain, the agent, and the host. These amounts can be determined experimentally by methods known in the art and as described above.

S. epidermidis infection treatment methods incorporate a therapeutically effective amount of an antibacterial agent or composition. A therapeutically effective amount is an amount sufficient to ameliorate or eliminate the infection. The prophylactically and/or therapeutically effective amounts can be administered in one administration or over repeated administrations. Therapeutic administration can be followed by prophylactic administration, once the initial bacterial infection has been resolved.

The antibacterial agents and compositions can be administered topically or systemically. Topical application is typically achieved by administration of creams, ointments, lotions, or sprays as described above. Systemic administration includes both oral and parental routes. Parental routes include, without limitation, subcutaneous, intramuscular, intraperitoneal, intravenous, transdermal, inhalation and intranasal administration.

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EXEMPLIFICATION

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Cloning and Sequencing S. epidermidis Genomic Sequence

This invention provides nucleotide sequences of the genome of *S. epidermidis* which thus comprises a DNA sequence library of *S. epidermidis* genomic DNA. The invention also provides nucleotide sequences of two naturally occurring plasmids in *S. epidermidis*. The detailed description that follows provides nucleotide sequences of *S. epidermidis*, and also describes how the sequences were obtained and how ORFs (Open Reading Frames) and protein-coding sequences can be identified. Also described are methods of using the disclosed *S. epidermidis* sequences in methods including diagnostic and therapeutic applications. Furthermore, the library can be used as a database for identification and comparison of medically important sequences in this and other strains of *S. epidermidis* as well as other species of *Staphylococcus*.

Chromosomal DNA from strain 18972 of *S. epidermidis*, was isolated using a protocol described by Storrs, et al.(*J. Bacteriol*. 173: 4347-4352 (1991). The only exception to this protocol was that lysostaphin (120 U/ml) was used instead of lysozyme. Two endogenous plasmids of approximately 39 Kb and 2.9 Kb in size were identified upon visualization of the *S. epidermidis* genomic DNA on a 0.5% agarose gel. The first library constructed contained fragments from the *S. epidermidis* genome as well as from the endogenous plasmid. A second library was later constructed with genomic DNA, from which the plasmid DNA was removed by CsCl centrifugation. The genomic DNA prep involved a lysozyme:lysostaphin digestion, sodium dodecyl sulfate lysis, Proteinase K and RNase treatment, phenol:chloroform extraction, and sodium acetate precipitation, followed by the CsCl gradient to remove the plasmid.

In the construction of both libraries, genomic *S. epidermidis* DNA was hydrodynamically sheared in an HPLC and then separated on a standard 1% agarose gel. A fraction corresponding to 2000-3000 bp in length was excised from the gel and purified by the GeneClean procedure (Bio101, Inc.).

The purified DNA fragments were then blunt-ended using T4 DNA polymerase. The healed DNA was then ligated to unique BstXI-linker adapters (5'-GTCTTCACCACGGGG-3' and 5'-GTGGTGAAGAC-3' in 100-1000 fold molar excess). These linkers are

complimentary to the BstXI-cut pGTC vector, while the overhang is not self-complimentary. Therefore, the linkers will not concatermerize nor will the cut-vector religate itself easily. The linker-adapted inserts were separated from the unincorporated linkers on a 1% agarose gel and purified using GeneClean. The linker-adapted inserts were then ligated to *BstXI*-cut vector to construct a "shotgun" sublclone libraries.

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Only major modifications to the protocols are highlighted. Briefly, the library was then transformed into DH5á competent cells (Gibco/BRL, DH5a transformation protocol). It was assessed by plating onto antibiotic plates containing ampicillin and IPTG/Xgal. The plates were incubated overnight at 37°C. Transformants were then used for plating of clones and picking for sequencing. The cultures were grown overnight at 37°C. DNA was purified using a silica bead DNA preparation (Engelstein, 1996) method. In this manner, 25 µg of DNA was obtained per clone.

These purified DNA samples were then sequenced using primarily ABI dye-terminator chemistry. All subsequent steps were based on sequencing by ABI377 automated DNA sequencing methods. The ABI dye terminator sequence reads were run on ABI377 machines and the data was transferred to UNIX machines following lane tracking of the gels. Base calls and quality scores were determined using the program PHRED (Ewing et al., 1998, Genome Res. 8: 175-185; Ewing and Green, 1998, Genome Res. 8: 685-734). Reads were assembled using PHRAP (P. Green, Abstracts of DOE Human Genome Program Contractor-Grantee Workshop V, Jan. 1996, p.157) with default program parameters and quality scores.

Finishing followed the initial assembly. Missing mates (sequences from clones that only gave reads from one end of the *Staphylococcus* DNA inserted in the plasmid) were identified and sequenced with ABI technology to allow the identification of additional overlapping contigs.

End-sequencing of randomly picked genomic lambda was also performed. Sequencing of both sides was done for all lambda sequences. The lambda library backbone helped to verify the integrity of the assembly and allowed closure of some of the physical gaps. Primers for walking off the ends of contigs would be selected using pick_primer (a GTC program) near the ends of the clones to facilitate gap closure. These walks can be sequenced using the selected clones and primers. These data are then reassembled with PHRAP. Additional

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sequencing using PCR-generated templates and screened and/or unscreened lambda templates can be done in addition.

Additional templates for the physical gaps were obtained through PCR using primers designed from the ends of the contigs. These templates were then used in sequencing reactions to close the gaps.

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Contigs were ordered by aligning identified *S. epidermidis* genes to the published physical maps. Order was confirmed by PCR. The final chromosomal assembly included 23 ordered contigs and the two plasmids each contained 1 contig each.

To identify *S. epidermidis* polypeptides the complete genomic sequence of *S.*10 epidermidis were analyzed essentially as follows: First, all possible stop-to- stop open reading frames (ORFs) greater than 180 nucleotides in all six reading frames were translated into amino acid sequences. Second, the identified ORFs were analyzed for homology to known (archeabacter, prokaryotic and eukaryotic) protein sequences. Third, the coding potential of non-homologous sequences were evaluated with the program GENEMARKTM (Borodovsky and McIninch, 1993, Comp. Chem. 17:123).

Identification, Cloning and Expression of S. epidermidis Nucleic Acids

Expression and purification of the *S. epidermidis* polypeptides of the invention can be performed essentially as outlined below.

To facilitate the cloning, expression and purification of membrane and secreted proteins from *S. epidermidis*, a gene expression system, such as the pET System (Novagen), for cloning and expression of recombinant proteins in *E. coli*, is selected. Also, a DNA sequence encoding a peptide tag, the His-Tag, is fused to the 3' end of DNA sequences of interest in order to facilitate purification of the recombinant protein products. The 3' end is selected for fusion in order to avoid alteration of any 5' terminal signal sequence.

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PCR Amplification and Cloning of Nucleic Acids Containing ORF's Encoding Enzymes

Nucleic acids chosen (for example, from the nucleic acids set forth in SEQ ID NO: 1-SEQ ID NO: 3772 for cloning from the 18972 strain of *S. epidermidis* and plasmids are prepared for amplification cloning by polymerase chain reaction (PCR). Synthetic oligonucleotide primers specific for the 5' and 3' ends of open reading frames (ORFs) are designed and purchased from GibcoBRL Life Technologies (Gaithersburg, MD, USA). All forward primers (specific for the 5' end of the sequence) are designed to include an NcoI cloning site at the extreme 5' terminus. These primers are designed to permit initiation of protein translation at a methionine residue followed by a valine residue and the coding sequence for the remainder of the native *S. epidermidis* DNA sequence. All reverse primers (specific for the 3' end of any *S. epidermidis* ORF) include a EcoRI site at the extreme 5' terminus to permit cloning of each *S. epidermidis* sequence into the reading frame of the pET-28b. The pET-28b vector provides sequence encoding an additional 20 carboxy-terminal amino acids including six histidine residues (at the extreme C-terminus), which comprise the His-Tag.

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Genomic DNA or plasmid DNA prepared from the 18972 strain of *S. epidermidis* is used as the source of template DNA for PCR amplification reactions (Current Protocols in Molecular Biology, John Wiley and Sons, Inc., F. Ausubel et al., eds., 1994). To amplify a DNA sequence containing an *S. epidermidis* ORF, genomic DNA (50 nanograms) is introduced into a reaction vial containing 2 mM MgCl₂, 1 micromolar synthetic oligonucleotide primers (forward and reverse primers) complementary to and flanking a defined *S. epidermidis* ORF, 0.2 mM of each deoxynucleotide triphosphate; dATP, dGTP, dCTP, dTTP and 2.5 units of heat stable DNA polymerase (Amplitaq, Roche Molecular Systems, Inc., Branchburg, NJ, USA) in a final volume of 100 microliters.

Upon completion of thermal cycling reactions, each sample of amplified DNA is washed and purified using the Qiaquick Spin PCR purification kit (Qiagen, Gaithersburg, MD, USA). All amplified DNA samples are subjected to digestion with the restriction

endonucleases, e.g., NcoI and EcoRI (New England BioLabs, Beverly, MA, USA)(Current Protocols in Molecular Biology, John Wiley and Sons, Inc., F. Ausubel et al., eds., 1994).

DNA samples are then subjected to electrophoresis on 1.0 % NuSeive (FMC BioProducts, Rockland, ME USA) agarose gels. DNA is visualized by exposure to ethidium bromide and long wave uv irradiation. DNA contained in slices isolated from the agarose gel is purified using the Bio 101 GeneClean Kit protocol (Bio 101 Vista, CA, USA).

Cloning of S. epidermidis Nucleic Acids Into an Expression Vector

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The pET-28b vector is prepared for cloning by digestion with restriction endonucleases, e.g., NcoI and EcoRI (Current Protocols in Molecular Biology, John Wiley and Sons, Inc., F. Ausubel et al., eds., 1994). The pET-28a vector, which encodes a His-Tag that can be fused to the 5¹ end of an inserted gene, is prepared by digestion with appropriate restriction endonucleases.

Following digestion, DNA inserts are cloned (Current Protocols in Molecular Biology, John Wiley and Sons, Inc., F. Ausubel et al., eds., 1994) into the previously digested pET-28b expression vector. Products of the ligation reaction are then used to transform the BL21 strain of *E. coli* (Current Protocols in Molecular Biology, John Wiley and Sons, Inc., F. Ausubel et al., eds., 1994) as described below.

20 Transformation Of Competent Bacteria With Recombinant Plasmids

Competent bacteria, *E coli* strain BL21 or *E. coli* strain BL21(DE3), are transformed with recombinant pET expression plasmids carrying the cloned *S. epidermidis* sequences according to standard methods (Current Protocols in Molecular, John Wiley and Sons, Inc., F. Ausubel et al., eds., 1994). Briefly, 1 microliter of ligation reaction is mixed with 50 microliters of electrocompetent cells and subjected to a high voltage pulse, after which, samples are incubated in 0.45 milliliters SOC medium (0.5% yeast extract, 2.0 % tryptone, 10 mM NaCl, 2.5 mM KCl, 10 mM MgCl2, 10 mM MgSO4 and 20, mM glucose) at 37 ©C with

shaking for 1 hour. Samples are then spread on LB agar plates containing 25 microgram/ml kanamycin sulfate for growth overnight. Transformed colonies of BL21 are then picked and analyzed to evaluate cloned inserts as described below.

5 Identification Of Recombinant Expression Vectors With S. epidermidis Nucleic Acids

Individual BL21 clones transformed with recombinant pET-28b *S. epidermidis* ORFs are analyzed by PCR amplification of the cloned inserts using the same forward and reverse primers, specific for each *S. epidermidis* sequence, that were used in the original PCR amplification cloning reactions. Successful amplification verifies the integration of the *S. epidermidis* sequences in the expression vector (Current Protocols in Molecular Biology, John Wiley and Sons, Inc., F. Ausubel et al., eds., 1994).

<u>Isolation and Preparation of Nucleic Acids From Transformants</u>

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Individual clones of recombinant pET-28b vectors carrying properly cloned *S*.

15 epidermidis ORFs are picked and incubated in 5 mls of LB broth plus 25 microgram/ml kanamycin sulfate overnight. The following day plasmid DNA is isolated and purified using the Qiagen plasmid purification protocol (Qiagen Inc., Chatsworth, CA, USA).

Expression Of Recombinant S. epidermidis Sequences In E. coli

The pET vector can be propagated in any *E. coli* K-12 strain e.g. HMS174, HB101, JM109, DH5, etc. for the purpose of cloning or plasmid preparation. Hosts for expression include *E. coli* strains containing a chromosomal copy of the gene for T7 RNA polymerase. These hosts are lysogens of bacteriophage DE3, a lambda derivative that carries the lacI gene, the lacUV5 promoter and the gene for T7 RNA polymerase. T7 RNA polymerase is induced by addition of isopropyl-B-D-thiogalactoside (IPTG), and the T7 RNA polymerase transcribes any target plasmid, such as pET-28b, carrying its gene of interest. Strains used include:

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BL21(DE3) (Studier, F.W., Rosenberg, A.H., Dunn, J.J., and Dubendorff, J.W. (1990) Meth. Enzymol. 185, 60-89).

To express recombinant *S. epidermidis* sequences, 50 nanograms of plasmid DNA isolated as described above is used to transform competent BL21(DE3) bacteria as described above (provided by Novagen as part of the pET expression system kit). The lacZ gene (betagalactosidase) is expressed in the pET-System as described for the *S. epidermidis* recombinant constructions. Transformed cells are cultured in SOC medium for 1 hour, and the culture is then plated on LB plates containing 25 micrograms/ml kanamycin sulfate. The following day, bacterial colonies are pooled and grown in LB medium containing kanamycin sulfate (25 micrograms/ml) to an optical density at 600 nM of 0.5 to 1.0 O.D. units, at which point, 1 millimolar IPTG was added to the culture for 3 hours to induce gene expression of the *S. epidermidis* recombinant DNA constructions.

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After induction of gene expression with IPTG, bacteria are pelleted by centrifugation in a Sorvall RC-3B centrifuge at 3500 x g for 15 minutes at 4 °C. Pellets are resuspended in 50 milliliters of cold 10 mM Tris-HCl, pH 8.0, 0.1 M NaCl and 0.1 mM EDTA (STE buffer). Cells are then centrifuged at 2000 x g for 20 min at 4 °C. Wet pellets are weighed and frozen at -80 °C until ready for protein purification.

A variety of methodologies known in the art can be utilized to purify the isolated proteins. (Current Protocols in Protein Science, John Wiley and Sons, Inc., J. E. Coligan et al., eds., 1995). For example, the frozen cells may be thawed, resupended in buffer and ruptured by several passages through a small volume microfluidizer (Model M-110S, Microfluidics International Corporation, Newton, MA). The resultant homogenate may be centrifuged to yield a clear supernatant (crude extract) and following filtration the crude extract may be fractionated over columns. Fractions may be monitored by absorbance at OD280 nm. and peak fractions may analyzed by SDS-PAGE

The concentrations of purified protein preparations may be quantified spectrophotometrically using absorbance coefficients calculated from amino acid content

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(Perkins, S.J. 1986 Eur. J. Biochem. 157, 169-180). Protein concentrations are also measured by the method of Bradford, M.M. (1976) Anal. Biochem. 72, 248-254, and Lowry, O.H., Rosebrough, N., Farr, A.L. & Randall, R.J. (1951) J. Biol. Chem. 193, pages 265-275, using bovine serum albumin as a standard.

SDS-polyacrylamide gels of various concentrations may be purchased from BioRad (Hercules, CA, USA), and stained with Coomassie blue. Molecular weight markers may include rabbit skeletal muscle myosin (200 kDa), *E. coli* (-galactosidase (116 kDa), rabbit muscle phosphorylase B (97.4 kDa), bovine serum albumin (66.2 kDa), ovalbumin (45 kDa), bovine carbonic anhydrase (31 kDa), soybean trypsin inhibitor (21.5 kDa), egg white

EQUIVALENTS

Those skilled in the art will recognize, or be able to ascertain using no more than routine experimentation, many equivalents to the specific embodiments and methods

15 described herein. The specific embodiments described herein are offered by way of example only, and the invention is to limited only by the terms of the appended claims, along with the full scope of equivalents to which such claims are entitled.

ORF Name	NTID	AAID NT AA score probability
AI7503000979_10192177_f1_14	1	3773 135 44
Description		
NO-HIT		
ORF Name	NTID	AAID LengthLength score probability
AI7503000979_10578392_f2_146	2	3774 510 169 525 1.7e-50
Description		
carrier proteindehydrase homolog (fa	hylococc bZ) gene precurso and Ten	us carnosus (3R)-hydroxymyristoyl acyl, partial cds, YwpF homolog, single-strand r (sceD), SceA precursor (sceA) and SceE A homolog (tenA) gene, partial cds.]
ORF Name	NTID	AAID <u>NT AA</u> LengthLength score probability
AI7503000979_10632763_f1_106	3	3775 135 44
Description		
NO-HIT	1.4	
ORF Name	NTID	AAID NT AA score probability
AI7503000979_10739063_c1_376	4	3776 693 230 601 1.5e-58
YwpF homolog, single-strand binding p	pept-bct roteinde rotein h (sceE) g	2] [DE:Staphylococcus carnosus hydrase homolog (fabZ) gene, partial cds, omolog (ssb), SceD precursor (sceD),SceA enes, completecds, and TenA homolog (tenA)
ORF Name	NTID	AAID <u>NT AA</u> score probability
AI7503000979_10939577_f1_79	5	3777 [153]50
Description NO-HIT		
ORF Name	NTID	AAID NT AA score probability
AI7503000979_1206255_f2_219	6	3778 126 41
Description		
NO-HIT		
ORF Name	NTID	AAID <u>NT AA</u> LengthLength
A17503000979_12111018_±3_297	7	3779 426 141 174 2.7e-13
Description		
<pre>gp:[GI:g642965] [LN:ABCARRA] [AC:X70 [DB:genpept-bct1] [DE:A.brasilense c</pre>		carR] [OR:Azospirillum brasilense] .] [NT:ORF2] [LE:59] [RE:580] [DI:direct]

ORF Name	NTID	AAID	NT AA LengthLength score probability
AT7503000979_12142768_f1_110	8	3780	300 99 110 4.2e-06
	>gp:[GI 466:AB00 R:Pyroco [DE:Pyro	:d103 9467:2 occus l	30234:g3256608] [LN:AP000001] :AB009468:AB009469] [PN:235aa long horikoshii] [SR:Pyrococcus horikoshii us horikoshii OT3 genomic DNA, 1-287000
ORF Name	NTID	AAID	NT AA score probability
AI7503000979_1250_c3_509	9	3781	
Description			
YwpF homolog, single-strand binding p precursor (sceA) and SceE precursor	pept-bct roteinde rotein h (sceE) g	2] [DI hydras nomolog genes,	DE:Staphylococcus carnosus ase homolog (fabZ) gene, partial cds,
ORF Name	NTID	AAID	NT AA score probability
AI7503000979_12587886_f1_33	10	3782	138 45
Description			
NO-HIT			
ORF Name	NTID		NT AA score probability
A17503000979_1281557_c3_517 Description	11	3783	1131 376 903 1.5e-90
sp:[LN:DDL_BACSU] [AC:P96612] [GN:DD:DD:DD:DD:DD:DD:DD:DD:DD:DD:DD:DD:DD	ALANYLAL D69613] ligase] 488] [AC d1A] [OR [EC:6.3. 35 and 5] [LN:B :dd1A] [2.4] [DE	ANINE [PN:I [OR:Ba ::AB001 ::Bacil 2.4] 47 dec SUB000 FN:per ::Bacil	C SYNTHETASE)] [SP:P96612] D-alanyl-D-alanine ligase A ddlA] sacillus subtilis] [DB:pir2] 1488] [PN:PROBABLE Illus subtilis] [SR:Bacillus subtilis [DE:Bacillus subtilis genome sequence, egree.] [LE:41311] [RE:42375] 103] [AC:Z99106:AL009126] 1011 ptidoglycan biosynthesis] [OR:Bacillus llus subtilis complete genome (section
ORF Name	NTID	AAID	NT AA score probability
AI7503000979_1292842_c2_415	12	3784	672 223 790 1.4e-78
Description		AD016:	4211 [OD Ghamballana
<pre>gp:[GI:d1037675:g4126674] [LN:AB01643 [SR:Staphylococcus aureus (strain:912 aureus, zinc responsible operon czr g</pre>	2) DNA]	[DB:ge	enpept-bct1] [DE:Staphylococcus
protein] [LE:2175] [RE:2813] [DI:dire			Tours parotar oab., [MI.Mypochecical

ORF Name	NTID	AAID	NT Length	<u>AA</u> Lengtl	score	proba	ability
AI7503000979_12_f1_1	13	3785	243	80	75	0.0084	
Description		L			J L	JL	
pir:[LN:A44803] [AC:A44803] [PN:pG1	proteir	n] [OR	:Homo	sapien	s] [SR	:, man]	[DB:pir2]
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Lengtl	score	proba	ability
AI7503000979_1366660_c3_484	14	3786	1215		566	7.8e-55	
Description				L	J L	J	
pir:[LN:A70842] [AC:A70842] [PN:proleuberculosis] [DB:pir2] >gp:[GI:e125] [PN:amiB] [GN:amiB] [OR:Mycobacteriumulosis H37Rv complete genome; 394. amiB, Probable] [LE:3363] [RE:4] [LN:MTV016] [AC:AL021841:AL123456] [DB:genpept] [DE:Mycobacteriumulosis tuberculosis H37Rv complete genome; 394. amiB, Probable] [LE:3363] [RE:4] [LN:MTV016] [AC:AL021841:AL123456] [DB:genpept] [DE:Mycobacteriumulosis tuberculosis [NT:Rv3306c, (MTV016.05c), len: 394. [DI:complement]	1137:g28 m tubero segment 547] [D] PN:amiB] culosis	394215 culosi 143/1 [:comp [GN:] [LN:1 s] [DB 62.] [1 lement] amiB] comple	MTV016 :genpej NT:Rv3: >gp: [OR:My6 ete gei	[AC: pt-bct 306c, [GI:e1 cobact nome;	AL021841 1] [DE:M (MTV016. 251137:g erium tu segment	l:AL123456] Mycobacterium .05c), len: g2894215] uberculosis]
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	proba	bility
AI7503000979_13790952_c2_422	15	3787	1194	397	1350	6.5e-13	8
Description							
[CL:phosphopentomutase] [OR:Bacillus [LN:BACJH642] [AC:D84432:D82370] [PN (strain:JH642(trpC2 PheA1)) DNA] [DB region containing skin element.] [LE >gp:[GI:e1185619:g2634785] [LN:BSUB00] [PN:phosphodeoxyribomutase] [GN:drm] [OR:Bacillus subtilis] [DB:genpept-bogenome (section 13 of 21): from 23952 [SP:P46353] [LE:51215] [RE:52399] [Dispense of the content of the c	:YqkN] :genpept :271220] 013] [AC [FN:cor ct1] [EC 261to 26	[OR:Bac c-bct1] [RE: C:Z991] nversic C:5.4.3	cillus] [DE:I 272404] 16:ALO on of 1 2.7] [I	subtil Bacillo [DI:0 09126] Cibose DE:Bac	lis] [us sub direct -1-P/d illus	SR:Bacil tilis DN] eoxyribo subtilis	llus subtilis NA, 283 Kb ose-1-P to] s complete
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	proba	bility
A17503000979_1415877_c2_413	16	3788	912	303		3.7e-16	
Description gp:[GI:g4982462] [LN:AE001824] [AC:AI protein] [GN:TM1876] [OR:Thermotoga r section 136 of 136 of the complete ge 60.17;] [LE:4717] [RE:5481] [DI:direct	maritima enome.]	l] [DB	genper	t-bct2	2] [DE	:Thermot	oga maritima
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	proba	bility
AI7503000979_14160455_c2_466	17	3789	162	53]		
Description							
NO-HIT	9.1- A	Tanket Law 1991			l velle <u>v</u> v		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	proba	bility
AI7503000979_14225327_f1_91	18	3790		342		1.6e-12	7
Description gp:[GI:d1037673:g4126672] [LN:AB01643 [OR:Staphylococcus aureus] [SR:Staphylococcus [DB:genpept-bct1] [DE:Staphylococcus completeand partial cds.] [NT:czcD]	lococcu aureus,	s aure zinc	eus (st respon	rain:9	12) Di operor	NA] n czr ge	-

ORF Name	LengthLength score probability
A17503000979_1438927_c1_356	19 3791 1686 561 2118 2.7e-219
SYNTHASE, (UTPAMMONIA LIGASE) (CTP >pir:[LN:SYBSTP] [AC:A32354:S55423:C6 synthase,:CTP-synthetase:UTPammonia [OR:Bacillus subtilis] [EC:6.3.4.2] [LN:BACSPOOFA] [AC:M22039] [PN:CTP syllate [SR:Bacillus subtilis (strains JH642 subtillis spoof, CTP synthetase (ctragenes, complete cds.] [LE:339] [RE:19 [AC:Z49782] [PN:CTP synthase] [GN:py:[DE:B.subtilis chromosomal DNA (region [RE:11087] [DI:direct] >gp:[GI:e11863 [PN:CTP synthetase] [GN:ctrA] [FN:py:[DB:genpept-bct1] [EC:6.3.4.2] [DE:Bacillus [EC:6.3.4.2]	TRA] [OR:BACILLUS SUBTILIS] [EC:6.3.4.2] [DE:CTP SYNTHETASE)] [SP:P13242] [DB:swissprot] 69610] [PN:CTP a ligase] [GN:ctrA:pyrG] [CL:CTP synthase] [DB:pirl] [MP:37 min] >gp:[GI:g143597] ynthetase] [GN:ctrA] [OR:Bacillus subtilis] and UOT0550) DNA] [DB:genpept-bctl] [DE:Bacillus A), andfructose-bisphosphate aldolase (orfY-tsr) 946] [DI:direct] >gp:[GI:g853762] [LN:BSDNA320D] rG] [OR:Bacillus subtilis] [DB:genpept-bctl] on 320-321 degrees).] [SP:P13242] [LE:9480] 216:g2636252] [LN:BSUB0020] [AC:Z99123:AL009126] rimidine biosynthesis] [OR:Bacillus subtilis] acillus subtilis complete genome (section 20 of ternate gene name: pyrG] [SP:P13242] [LE:11346]
ORF Name AT7503000979_14454660_f1_26 Description NO-HIT	NTID AAID NT AA score probability 20 3792 138 45
ORF Name AI7503000979_14492142_f1_81 Description NO-HIT	NTID AAID NT AA score probability 21 3793 126 41
<pre>prowazekii] [DB:pir2] >gp:[GI:e134259 [PN:unknown] [GN:RP285] [OR:Ricketts]</pre>	NTID AAID NT AA LengthLength score probability 22 3794 171 56 52 0.029 othetical protein RP285] [GN:RP285] [OR:Rickettsia 90:g3860846] [LN:RPXX02] [AC:AJ235271:AJ235269] ia prowazekii] [DB:genpept-bct1] [DE:Rickettsia genome; segment2/4.] [LE:68773] [RE:69213]
[OR:Pyrococcus horikoshii] [DB:pir2]	NTID AAID NT AA Score probability 23 3795 294 97 133 6.0e-09 othetical protein PHS004] [GN:PHS004] >gp:[GI:d1030236:g3256610] [LN:AP000001] 466:AB009467:AB009468:AB009469] [PN:58aa long

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
AI7503000979_14855051_c1_401	24	3796	471	156	409	3.4e-38
Description pir: [LN:H69773] [AC:H69773] [PN:con [CL:hypothetical protein HI1173] [OR >gp: [GI:d1020070:g1881290] [LN:AB001 [SR:Bacillus subtilis (strain:168) D sequence, 148 kb sequence of the reg UNKNOWN.] [LE:61365] [RE:61817] [DI: [AC:Z99106:AL009126] [GN:ydcK] [FN:u [DE:Bacillus subtilis complete genom [NT:similar to hypothetical proteins ORF Name	served harmonic served harmoni	nypothus subgenperson 35 sqp:[GR:Basen 3 Griden	etical tilis] 1488] pt-bctl and 47 GI:ell8 acillus of 21):	protei [DB:pi [GN:ydc 1] [DE: 7 degre 32445:g 5 subti : from 25562]	n ydch r2] :K] [OH Bacil] :e.] [N :263277 lis] 402751	GN:ydcK] R:Bacillus subtilis] Lus subtilis genome NT:FUNCTION [IN:BSUB0003] [DB:genpept-bct1] L to611850.]
A17503000979_15798901_c3_471 Description NO-HIT	25	3797		40		
ORF Name A17503000979_16251305_c3_501 Description sp:[LN:ATP6_BACST] [AC:P42010] [GN:A [DE:ATP SYNTHASE A CHAIN, (PROTEIN 6 >gp:[GI:d1007828:g534857] [LN:BACATP stearothermophilus] [SR:Bacillus ste o] [DB:genpept-bct1] [DE:Bacillus st c,complete cds.] [LE:256] [RE:966] [)] [SP:P SAC] [AC arotherm earother	42010 ::D380! ophilu mophil	LLUS ST] [DB:s 59] [PN us (str	ZEAROTH Swisspr I:ATPas	ERMOPH ot] e subu	nit a] [OR:Bacillus (library: library
ORF Name A17503000979_165888_f3_321 Description NO-HIT	NTID	ΔΔΤΩ	NT Length	AA Length 51	score	probability
ORF Name A17503000979_19693831_c1_350 Description NO-HIT		<u>AAID</u> 3800	NT Length	AA Length 54	score	probability
ORF Name AT7503000979_19728433_f2_133 Description sp:[LN:YWCF_BACSU] [AC:P39604] [GN:YI [DE:HYPOTHETICAL 43.3 KD PROTEIN IN (IDE:Swissprot] >pir:[LN:S39697] [AC:YWCF:protein ipa-42d] [GN:YWCF] [OR [LN:BSGENR] [AC:X73124] [GN:ipa-42d] [DE:B.subtilis genomic region (325 to [DI:direct] >gp:[GI:e1186311:g263634' [FN:unknown] [OR:Bacillus subtilis] genome (section 20 of 21): from 37984	WCF:IPA- QOXD-VPR S39697:A :Bacillu [OR:Bac 0 333).] 7] [LN:B	42D] INTER 70053 s subt illus [SP:F SUB002 ept-bo	RGENIC [PN: cilis] subtil P39604] 20] [AC	ILLUS REGION Cell-d: [DB:pi: is] [DI [LE:44:29912]	SUBTIL] [SP: ivisio r2] >g B:genp 4053] 3:AL00 llus s	P39604] n protein homolog p:[GI:g413966] ept-bct1] [RE:45234] 9126] [GN:ywcF] ubtilis complete
similar to] [SP:P39604] [LE:112728]	[RE:1139	09] [[I:comp	lement]	

ORF Name NTID AAID LengthLength score probability

A17503000979_19739675_c3_494 30 3802 636 211 638 1.8e-62

Description

sp:[LN:KITH BACSU] [AC:Q03221] [GN:TDK] [OR:BACILLUS SUBTILIS] [EC:2.7.1.21] [DE:THYMIDINE KINASE,] [SP:Q03221] [DB:swissprot] >pir:[LN:S55432] [AC:S55432:D69721] [PN:thymidine kinase, tdk] [GN:tdk] [CL:thymidine kinase] [OR:Bacillus subtilis] [EC:2.7.1.21] [DB:pir2] >gp:[GI:g405819] [LN:BACRHOTDKX] [AC:M97678] [PN:thymidine kinase] [GN:tdk] [OR:Bacillus subtilis] [SR:Bacillus subtilis (Transposon Tn917 insertional library) DNA] [DB:genpept-bct1] [DE:Bacillus subtilis OrfR, 3' end; OrfO; transcriptional terminator(rho) gene; ribosomal protein L31; thymidine kinase (tdk) gene, complete cds.] [NT:Incorrect sequence given in Quirk et al. citation,] [LE:3334] [RE:3921] [DI:direct] >qp:[GI:q853771] [LN:BSDNA320D] [AC:Z49782] [PN:thymidine kinase] [GN:tdk] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:B.subtilis chromosomal DNA (region 320-321 degrees).] [SP:Q03221] [LE:18786] [RE:19373] [DI:direct] >gp:[GI:e1184612:g2636231] [LN:BSUB0019] [AC:Z99122:AL009126] [PN:thymidine kinase] [GN:tdk] [OR:Bacillus subtilis] [DB:genpept-bct1] [EC:2.7.1.21] [DE:Bacillus subtilis complete genome (section 19 of 21): from 3597091to 3809700.] [SP:Q03221] [LE:204370] [RE:204957] [DI:complement] >gp:[GI:e1186207:g2636243] [LN:BSUB0020] [AC:Z99123:AL009126] [PN:thymidine kinase] [GN:tdk] [OR:Bacillus subtilis] [DB:genpept-bct1] [EC:2.7.1.21] [DE:Bacillus subtilis complete genome (section 20 of 21): from 3798401to 4010550.] [SP:Q03221] [LE:3060] [RE:3647] [DI:complement] >gp:[GI:e1184612:g2636231] [LN:BSUB0019] [AC:Z99122:AL009126] [PN:thymidine kinase] [GN:tdk] [OR:Bacillus subtilis] [DB:genpept] [EC:2.7.1.21] [DE:Bacillus subtilis complete genome (section 19 of 21): from 3597091to 3809700.] [SP:Q03221] [LE:204370] [RE:204957] [DI:complement]

ORF Name	NTID	AAID <u>NT AA</u> LengthLength score probability
A17503000979_20032527_f2_127	31	3803 165 54
Description NO-HIT		
ORF Name	NTID	AAID NT AA score probability
AI7503000979_20156686_c3_498	32	3804 573 190 415 7.8e-39

Description

sp:[LN:YWLG_BACSU] [AC:P39157] [GN:YWLG:IPC-33D] [OR:BACILLUS SUBTILIS]
[DE:HYPOTHETICAL 19.4 KD PROTEIN IN SPOIIR-GLYC INTERGENIC REGION] [SP:P39157]
[DB:swissprot] >pir:[LN:I40482] [AC:I40482:D70062:S49362] [PN:hypothetical protein
ywlG:ipc-33d protein] [GN:ywlG:ipc-33d] [OR:Bacillus subtilis] [DB:pir2]
>gp:[GI:g556885] [LN:BSSPORUPP] [AC:Z38002] [PN:Unknown] [GN:ipc-33d] [OR:Bacillus
subtilis] [DB:genpept-bct1] [DE:B.subtilis spoII-R, glyC and upp genes.] [SP:P39157]
[LE:4750] [RE:5292] [DI:direct] >gp:[GI:e1184597:g2636216] [LN:BSUB0019]
[AC:Z99122:AL009126] [GN:ywlG] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1]
[DE:Bacillus subtilis complete genome (section 19 of 21): from 3597091to 3809700.]
[NT:alternate gene name: ipc-33d] [SP:P39157] [LE:192609] [RE:193151] [DI:complement]
>gp:[GI:e1184597:g2636216] [LN:BSUB0019] [AC:Z99122:AL009126] [GN:ywlG] [FN:unknown]
[OR:Bacillus subtilis] [DB:genpept] [DE:Bacillus subtilis complete genome (section 19 of 21): from 3597091to 3809700.] [NT:alternate gene name: ipc-33d] [SP:P39157]
[LE:192609] [RE:193151] [DI:complement]

ORF Name	NTID	AAID LengthLength score probability
AI7503000979_20348453_c3_518	33	3805 1587 528 1226 9.0e-125
[GN:aggH] [OR:Lactobacillus reuteri]	[DB:gei ggH)gen	PN:autoaggregation-mediating protein enpept-bct2] [DE:Lactobacillus reutering e, complete cds.] [NT:AggH; putative e] [RE:1674] [DI:direct]
ORF Name AI7503000979_20360687_c3_516 Description NO-HIT	NTID 34	AAID NT AA score probability 3806 141 46
ORF Name AT7503000979_20569052_c2_457	NTID	AAID NT AA score probability LengthLength 198 7.8e-16
Description		250 1
pir:[LN:G70041] [AC:G70041] [PN:con [OR:Bacillus subtilis] [DB:pir2] >gp [AC:Z99121:AL009126] [GN:yvgZ] [FN:u	:[GI:e1: nknown] e (sect:	[OR:Bacillus subtilis] [DB:genpept-bct1] ion 18 of 21): from 3399551to 3609060.]
ORF Name	NTID	AAID NT AA score probability
AI7503000979_20572255_f1_34	36	3808 207 68
Description NO-HIT		
ORF Name AI7503000979_20900062_f2_215	NTID	AAID NT AA score probability 3809 147 48 53 0.045
Description		
	istol Na ains wea	
ORF Name	NTID	AAID NT AA score probability
A17503000979_2117125_c2_432	38	3810 129 42
Description NO NATE		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000979_2125637_£3_335	39	3811 150 49
Description		
NO-HIT		

ORF Name	NTID	AAID LengthLength score probability
A17503000979_21517182_±3_322	40	3812 375 124 396 8.1e-37
Description		
completeand partial cds.] [LE:1703] [LN:AF044951] [AC:AF044951] [PN:reprtransport repressor] [OR:Staphylocod	ylococc aureus [RE:202] essor p	us aureus (strain:912) DNA] , zinc responsible operon czr genes,
ORF Name	NTID	AAID NT AA score probability
A17503000979_21562827_£3_306	41	3813 132 43
Description	L	
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000979_21756937_c1_379	42	3814 672 223 401 2.4e-37
Description		
pyrophosphorylase, thiC:protein ipa- pyrophosphorylase homology] [OR:Baci >gp:[GI:g413950] [LN:BSGENR] [AC:X73 [DB:genpept-bct1] [DE:B.subtilis gen [RE:26856] [DI:direct] >gp:[GI:e1186 [PN:thiamine-phosphate pyrophosphory pyrophosphate of] [OR:Bacillus subti	(TMP-PPA \$39681] 26d] [GI 1lus sul 124] [GI comic rec 328:g263 (lase] [CI lof 21)	ASE) (THIAMIN-PHOSPHATE SYNTHASE)] [AC:S39681:E69722] [PN:thiamin-phosphate N:thiC] [CL:thiE protein:thiamin-phosphate btilis] [EC:2.5.1.3] [DB:pir2] N:ipa-26d] [OR:Bacillus subtilis] gion (325 to 333).] [SP:P39594] [LE:26188] 36364] [LN:BSUB0020] [AC:Z99123:AL009126] GN:thiC] [FN:substitution of the B:genpept-bct1] [EC:2.5.1.3] [DE:Bacillus : from 3798401to 4010550.] [NT:alternate
ORF Name	NTID	AAID LengthLength score probability
A17503000979_2230303_f3_265	43	3815 669 222 346 1.6e-31
KD PROTEIN IN BSAA-ILVD INTERGENIC R [AC:E69935] [PN:conserved hypothetical protein AF0994] [OR:Bace [LN:BACYACA] [AC:L77246] [GN:ypgQ] [[DE:Bacillus subtilis (YAC10-9 clone [NT:putative] [LE:16914] [RE:17531] [LN:BSUB0012] [AC:Z99115:AL009126] [[DB:genpept-bct1] [DE:Bacillus subtilis]	EGION] cal prot illus su OR:Bacil) DNA re [DI:dire GN:ypgQl lis comp	ubtilis] [DB:pir2] >gp:[GI:g1256633] llus subtilis] [DB:genpept-bct1] egion between the serA andkdg loci.]

[RE:108238] [DI:complement]

ORF Name	NTID	AAID LengthLength score probability
AI7503000979_22460882_c3_482	44	3816 687 228 612 1.0e-59
Description		
	genpept	82174] [PN:deoxyribose-phosphate aldolase] -bct1] [EC:4.1.2.4] [DE:B.subtilis operon] [LE:1462] [RE:2106] [DI:direct]
ORF Name	NTID	AAID NT AA score probability
A17503000979_22692137_c2_464	45	3817 486 161 688 9.2e-68
Description		
aureus] [DB:genpept-bct1] [DE:S.aure >gp:[GI:e284999:g1729796] [LN:SAUSIG	us sigB B] [AC:	:Y07645] [GN:rsbW] [OR:Staphylococcus gene.] [LE:2220] [RE:2699] [DI:direct] [Y09929] [GN:rsbW] [OR:Staphylococcus aureus] [rsbW & sigB genes.] [LE:2233] [RE:2712]
ORF Name	NTID	AAID NT AA score probability
AI7503000979_23437803_c3_523	46	Length Length Score Disability
Description		
		:Y07645] [GN:ORF6] [OR:Staphylococcus gene.] [LE:3878] [RE:>5272] [DI:direct]
ORF Name	NTID	AAID NT AA score probability
AI7503000979_23439002_c2_417	47	3819 1404 467
Description NO-HIT		
ORF Name	NTID	AAID NT AA score probability
AI7503000979_23446887_f3_323	48	3820 1155 384 548 6.3e-53
Description		
<pre>gp:[GI:g4097757] [LN:SAU67965] [AC:U [OR:Staphylococcus aureus] [DB:genpe protein gene, complete cds.] [LE:712</pre>	pt-bct2	[DE:Staphylococcus aureus lytic regulatory
ORF Name	NTID	AAID NT AA score probability
AI7503000979_23594057_c1_362	49	
Description		
NO-HIT		

ORF Name	NTID	AAID LengthLength score probability					
A17503000979_23595137_c3_480	50	3822 501 166 423 1.1e-39					
sigma-B dps] [GN:dps] [CL:hypothet [DB:pir2] >gp:[GI:e1185938:g2635549] [OR:Bacillus subtilis] [DB:genpept-Left of 21): from 2997771to 3213410.] [SP:P80879] [LE:137548] [RE:137985] [AC:AF008220] [GN:ytkB] [OR:Bacillus	ical pro [LN:BSU pctl] [DI [NT:alte [DI:comp subtili	UB0016] [AC:Z99119:AL009126] [GN:dps] E:Bacillus subtilis complete genome (section ernate gene name: ytkB; stress- and] plement] >gp:[GI:g2293159] [LN:AF008220] is] [DB:genpept-bct2] [DE:Bacillus subtilis o HI1349 from H. influenzae] [LE:42442]					
ORF Name	NTID	AAID LengthLength score probability					
A17503000979_23625008_c1_347	51	3823 963 320 749 3.2e-74					
DNA, clone_lib:lambda no.] [DB:genpe	lodurans pt-bct1] .C004.]	:AB011838] [PN:mannnose-6 phospate s] [SR:Bacillus halodurans (strain:C-125) [EC:5.3.1.8] [DE:Bacillus halodurans C-125 [NT:similar to B.subtilis ydhS gene(53-69%					
ORF Name	NTID	AAID NT AA score probability					
A17503000979_23625387_c1_402	52	3824 219 72					
NO-HIT							
ORF Name	NTID	AAID NT AA score probability					
A17503000979_23634678_c2_424	53	3825 1194 397					
Description NO-HIT							
ORF Name	NTID	AAID NT AA score probability					
A17503000979_23634702_c2_453	54	3826 792 263 498 1.3e-47					
Description sp:[LN:THIM_BACSU] [AC:P39593] [GN:THIM:THIK:IPA-25D] [OR:BACILLUS SUBTILIS] [EC:2.7.1.50] [DE:HYDROXYETHYLTHIAZOLE KINASE) (THZ KINASE) (TH KINASE)] [SP:P39593] [DB:swissprot] >pir:[LN:S39680] [AC:S39680:G69722] [PN:hydroxyethylthiazole kinase, thiK:protein ipa-25d] [GN:thiK] [CL:hydroxyethylthiazole kinase: hydroxyethylthiazole kinase homology] [OR:Bacillus subtilis] [EC:2.7.1.50] [DB:pir2] >gp:[GI:g413949] [LN:BSGENR] [AC:X73124] [GN:ipa-25d] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:B.subtilis genomic region (325 to 333).] [SP:P39593] [LE:25373] [RE:26191] [DI:direct] >gp:[GI:e1186329:g2636365] [LN:BSUB0020] [AC:Z99123:AL009126] [PN:hydroxyethylthiazole kinase] [GN:thiK] [FN:phosphorylation of] [OR:Bacillus subtilis] [DB:genpept-bct1] [EC:2.7.1.50] [DE:Bacillus subtilis complete genome (section 20 of 21): from 3798401to 4010550.] [NT:alternate gene name: ipa-25d, ywbJ] [SP:P39593] [LE:131768] [RE:132586] [DI:complement]							

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000979_23651702_c1_343	55	3827	870	289	255	7.1e-22
Description pir: [LN:C70070] [AC:C70070] [PN:con [CL:hypothetical protein ywpJ] [OR:B >gp: [GI:e1184491:g2636110] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-b 19 of 21): from 3597091to 3809700.] [RE:98190] [DI:direct] >gp: [GI:e3080 [OR:Bacillus subtilis] [DB:genpept-b genes.] [NT:product similar to Bacil [DI:complement] >gp: [GI:e1184491:g26 [FN:unknown] [OR:Bacillus subtilis] (section 19 of 21): from 3597091to 3 [LE:97330] [RE:98190] [DI:direct]	acillus 019] [A ct1] [D [NT:sim 93:g189 ct1] [D lus sub 36110] [DB:gen	subti C:Z991 E:Baci ilar t 4770] E:B.su tilis [LN:BS pept]	lis] [1 22:AL0 1lus s o hypo [LN:BS btilis YxeH a UB0019 [DE:Ba	DB:pir2 09126] ubtilis thetica Z92954] yws[A, nd YcsE] [AC:Z	[GN:yv comp] l prot [AC:2 B,C,D,] [LE: 99122: subtil	vtE] [FN:unknown] Lete genome (section ceins] [LE:97330] LE:9754] [GN:ywtE] LE:F,G] and gerBC LE:5152] LE:AL009126] [GN:ywtE] Lis complete genome
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
AI7503000979_23711642_c1_397	56	3828	1026			5.6e-148
Description gp:[GI:e284997:g1729794] [LN:SAUSIGB [DB:genpept-bct1] [DE:S.aureus rsbU, [DI:direct]						
ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	score	probability
A17503000979_2379658_f2_237	57	3829	225	74		
Description NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000979_2383253_c2_405	58	3830	399	132		
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
AI7503000979_24015687_c2_438	59	3831	465	154	119	1.8e-07
Description sp:[LN:ATPZ_BACP3] [AC:P09354] [OR:BI [DE:ATP SYNTHASE PROTEIN I] [SP:P093] [PN:H+-transporting ATP synthase, chi chain I] [OR:thermophilic bacterium I [LN:PS3TF0F1] [AC:X07804:X07374] [OR [DE:Thermophilic bacterium PS3 TF0F-1 (AA 1 - 127)] [SP:P09354] [LE:433] [I ORE Name	54] [DB ain I] PS-3] [1 :thermon 1 operon RE:816]	:swiss [CL:Bac EC:3.6 philic n for A [DI:d:	prot] : cillus .1.34] bacter ATP syr	Ppir:[Li H+-tran [DB:pi: rium PS: nthase (N:S013 nsport r2] >g 3] [DB comple	97] [AC:S01397] ing ATP synthase p:[GI:g45809] :genpept-bct1] x.] [NT:I protein
ORF Name	NTID	_	Length	Length	score	probability
A17503000979_24074137_c3_492	60	3832	1332	443	1360	5.7e-139
Description gp:[GI:g143434] [LN:BACRHOTDKX] [AC:I [FN:transcriptional terminator] [OR:I (Transposon Tn917 insertional library OrfR, 3' end; OrfQ; transcriptional thymidine kinase (tdk) gene, complete	Bacillus y) DNA] terminat	s subti DB:ge tor(rhe	ilis] (enpept- o) gene	[SR:Bac: -bct1] e; ribos	illus [DE:Ba somal	cillus subtilis protein L31;

ORF Name	NTID	AAID	Length Length score probability
A17503000979_24228411_c1_375	61	3833	
Description		J	
gp:[GI:g2735512] [LN:SCU96108] [AC:U [GN:ssb] [OR:Staphylococcus carnosus (3R)-hydroxymyristoyl acyl carrier p YwpF homolog, single-strand binding p precursor (sceA) and SceE precursor gene, partial cds.] [NT:SSB] [LE:857] [DB:qroteing roteing (sceE)	genpept dehydra homolo genes,	-bct2] [DE:Staphylococcus carnosus se homolog (fabZ) gene, partial cds, g (ssb), SceD precursor (sceD),SceA completecds, and TenA homolog (tenA)
ORF Name	NTID	AAID	NT AA LengthLength score probability
AI7503000979_24235952_c1_398 Description	62	3834	327 108 501 6.0e-48
			[GN:rsbV] [OR:Staphylococcus aureus] sigB genes.] [LE:1905] [RE:2231]
ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000979_24245327_c1_364	63	3835	714 237 843 3.5e-84
Description			
<pre>gp:[GI:e258331:g1765902] [LN:BCUPPGL phosphoribosyltransferase] [GN:upp] [DE:B.caldolyticus upp gene.] [SP:P7</pre>	[OR:Bac	cillus (caldolyticus] [DB:genpept-bct1]
ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000979_24254202_c1_368	64	3836	1512 503 2050 4.3e-212
Description sp:[LN:ATPA_BACME] [AC:P17674] [GN:A' SYNTHASE ALPHA CHAIN,] [SP:P17674] [IPN:H+-transporting ATP synthase, all chain: H+-transporting ATP synthase [EC:3.6.1.34] [DB:pir2] >gp:[GI:g142] [PN:ATP synthase alpha subunit] [OR:DNA, clones pWSB100, pCAH1.3, and pW synthase i,a,c,b,delta,alpha,gamma,boorf.] [LE:2853] [RE:4361] [DI:direct	DB:swis pha cha alpha c 559] [I Bacillu PC208] eta anc	ssprot] ain] [C] chain ho LN:BACA' us megan [DB:gen	>pir:[LN:F31482] [AC:F31482] L:H+-transporting ATP synthase alpha bmology] [OR:Bacillus megaterium] TPA] [AC:M20255:J04455:M18352:M23924] terium] [SR:B.megaterium (QM B1551) hpept-bct1] [DE:B.megaterium ATP
ORF Name	NTID	AAID	NT AA score probability
A17503000979_24256562_c1_381	65	3837	126 41
Description NO-HIT			
ORF Name	NTID	AAID	<u>NT AA</u> LengthLength
AI7503000979_24350953_f1_2	66	3838	132 43 72 0.017
Description			
	>gp:[0 466:AB0 R:Pyroc [DE:Pyr	31:d1030 009467: <i>l</i> coccus l	D229:g3256603] [LN:AP000001] AB009468:AB009469] [PN:106aa long Aorikoshii] [SR:Pyrococcus horikoshii s horikoshii OT3 genomic DNA, 1-287000

ORF Name	NTID	AAID NT AA score probability
AI7503000979_24353427_c3_503	67	3839 543 180 229 4.0e-19
SYNTHASE DELTA CHAIN,] [SP:P37811] [[AC:I40365:D69592:S39253] [PN:H+-tr [GN:atpH] [CL:H+-transporting ATP s [EC:3.6.1.34] [DB:pir2] >gp:[GI:g433 subunit delta] [GN:atpH] [OR:Bacillu atpase genes for ATP synthase subuni [SP:P37811] [LE:2484] [RE:3029] [DI: [AC:Z99122:AL009126] [PN:ATP synthas [DB:genpept-bct1] [EC:3.6.1.34] [DE: 21): from 3597091to 3809700.] [SP:P3 >gp:[GI:e1184590:g2636209] [LN:BSUB0 delta)] [GN:atpH] [OR:Bacillus subti	DB:swissansport: ynthase 988] [Lissing subtile ts i, agoing direct] e (subur) Bacillus 7811] [1] 019] [AGoing] [DIO of 21)	OR:BACILLUS SUBTILIS] [EC:3.6.1.34] [DE:ATP sprot] >pir:[LN:I40365] ting ATP synthase, delta chain (atpH)] e delta chain] [OR:Bacillus subtilis] LN:BSATPASE] [AC:Z28592] [PN:ATP synthase ilis] [DB:genpept-bct1] [DE:B.subtilis (168) a, c ,b, delta, alpha, gamma, beta, epsilon.] >pp:[GI:e1184590:g2636209] [LN:BSUB0019] unit delta)] [GN:atpH] [OR:Bacillus subtilis] us subtilis complete genome (section 19 of [LE:187368] [RE:187913] [DI:complement] AC:Z99122:AL009126] [PN:ATP synthase (subunit DB:genpept] [EC:3.6.1.34] [DE:Bacillus o: from 3597091to 3809700.] [SP:P37811]
ORF Name AI7503000979_24392193_c1_346	NTID 68	AAID NT AA score probability 3840 204 67
Description		l [][]
NO-HIT		No.
ORF Name	NTID	AAID NT AA score probability
>pir:[LN:G32354] [AC:S55428:G32354:H 1-carboxyvinyltransferase, murZ] [GN 1-carboxyvinyltransferase MurZ] [OR: >gp:[GI:g853767] [LN:BSDNA320D] [AC: [OR:Bacillus subtilis] [DB:genpept-b degrees).] [SP:P19670] [LE:14403] [R [LN:BSUB0019] [AC:Z99122:AL009126] [[FN:peptidoglycan biosynthesis] [OR: [DE:Bacillus subtilis complete genom [NT:alternate gene name: murZ, lssf, >gp:[GI:e1186211:g2636247] [LN:BSUB0 [PN:UDP-N-acetylglucosamine] [GN:mur subtilis] [DB:genpept-bct1] [EC:2.5. 20 of 21): from 3798401to 4010550.] [LE:6741] [RE:8030] [DI:complement] [AC:Z99122:AL009126] [PN:UDP-N-acetylbiosynthesis] [OR:Bacillus subtilis]	RASE) (FRASE)	(EPT)] [SP:P19670:Q03225] [DB:swissprot] [PN:UDP-N-acetylglucosamine [CL:UDP-N-acetylglucosamine as subtilis] [EC: 2.5.1.7] [DB:pir2] [PN:UDP-N-acetylglucosamine] [GN:murZ] [E:B.subtilis chromosomal DNA (region 320-321) [C] [DI:direct] >gp:[GI:e1184616:g2636235] [N-acetylglucosamine] [GN:murZ] [IS subtilis] [DB:genpept-bct1] [EC:2.5.1.7] [IS subtilis] [DB:genpept-bct1] [EC:2.5.1.7] [IO 19 of 21): from 3597091to 3809700.] [LE:208051] [RE:209340] [DI:complement] [IC:299123:AL009126] [IS peptidoglycan biosynthesis] [OR:Bacillus [IS Bacillus subtilis complete genome (section cernate gene name: murZ, lssF, rev-4] [II 184616:g2636235] [LN:BSUB0019] [II 184616:g2636235] [IN:BSUB0019] [II 184616:g2636235] [IN:BSUB0019]
ORF Name A17503000979_24407631_t3_252	NTID	AAID NT AA score probability 3842 144 47
Description		

NO-HIT

NT ORF Name score probability LengthLength A17503000979_24415933_c2_418 3843 144 435 0.026 Description sp:[LN:YUXK_BACSU] [AC:P40761:005233] [GN:YUXK] [OR:BACILLUS SUBTILIS] [DE:HYPOTHETICAL 15.7 KD PROTEIN IN PBPD-COMA INTERGENIC REGION (ORF2)] [SP:P40761:005233] [DB:swissprot] >pir:[LN:B55220] [AC:B55220:D70025] [PN:hypothetical protein yuxK:pbpD 3'-region hypothetical protein] [GN:yuxK] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:g520537] [LN:BSU11882] [AC:U11882] [PN:unknown] [GN:pbpD] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis orf1, partial cds, penicillin-binding protein 4(pbpD) gene, complete cds, and orf2, complete cds.] [NT:orf2] [LE:2760] [RE:3173] [DI:direct] >qp:[GI:e1184228:q2635646] [LN:BSUB0017] [AC:Z99120:AL009126] [GN:yuxK] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 17 of 21): from 3197001to 3414420.] [NT:alternate gene name: yugD] [SP:P40761] [LE:37885] [RE:38298] [DI:direct] >gp:[GI:e311522:q1934785] [LN:BSZ93933] [AC:Z93933] [PN:unknown] [GN:yugD] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:B.subtilis genomic DNA fragment from yugA to yugD.] [SP:P40761] [LE:7318] [RE:7731] [DI:direct] NT AΑ ORF Name NTID <u>score</u> AAID probability LengthLength A17503000979_24508563_c2_436 3844 1092 363 9.0e-125 Description sp:[LN:RF1 BACSU] [AC:P45872] [GN:PRFA] [OR:BACILLUS SUBTILIS] [DE:PEPTIDE CHAIN RELEASE FACTOR 1 (RF-1)] [SP:P45872] [DB:swissprot] >pir:[LN:S55437] [AC:S55437:G69681] [PN:translation releasing factor RF-1:peptide chain release factor 1] [GN:prfA:RF-1] [CL:translation releasing factor] [OR:Bacillus subtilis] [DB:pir1] >gp:[GI:g853776] [LN:BSDNA320D] [AC:Z49782] [PN:peptide chain release factor 1] [GN:prfA] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:B.subtilis chromosomal DNA (region 320-321 degrees).] [NT:gtg start codon] [SP:P45872] [LE:23623] [RE:24693] [DI:direct] >gp:[GI:e1184607:g2636226] [LN:BSUB0019] [AC:Z99122:AL009126] [PN:peptide chain release factor 1] [GN:prfA] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 19 of 21): from 3597091to 3809700.] [SP:P45872] [LE:199050]

[RE:200120] [DI:complement] >gp:[GI:e1184607:g2636226] [LN:BSUB0019]

ORF Name

Description NO-HIT

A17503000979 24625216 f3 264

3597091to 3809700.] [SP:P45872] [LE:199050] [RE:200120] [DI:complement]

[AC:Z99122:AL009126] [PN:peptide chain release factor 1] [GN:prfA] [OR:Bacillus

NTID

subtilis] [DB:genpept] [DE:Bacillus subtilis complete genome (section 19 of 21): from

AAID

3,845

NT

186

LengthLength

AA

<u>score</u>

probability

ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000979_24641687_c2_437	74	3846	
Description	/ -	5010	
sp:[LN:YWLE_BACSU] [AC:P39155] [GN:YI [DE:(EC 3.1.3.48)] [SP:P39155] [DB:si] [PN:protein-tyrosine-phosphatase,] [GN:ywlE:ipc-31d] [CL:protein-tyrosisubtilis] [EC:3.1.3.48] [DB:pir1] >gp [PN:Unknown] [GN:ipc-31d] [OR:BacillisspoII-R, glyC and upp genes.] [SP:P39] >gp:[GI:el184599:g2636218] [LN:BSUB00]	wissprot homolog ine-phos p:[GI:g5 us subti 9155] [I 019] [AC ct1] [DE [NT:alte [DI:comp GN:ywlE] omplete	ywlE, sphata 56883 lis] E:367 C:2991 C:Baci ernate clemen [FN: genom	r:[LN:S49360] [AC:I40479:B70062:S49360 low molecular weight:ipc-31d protein] se, low molecular weight] [OR:Bacillus] [LN:BSSPORUPP] [AC:Z38002] [DB:genpept-bct1] [DE:B.subtilis 9] [RE:4131] [DI:direct] 22:AL009126] [GN:ywlE] [FN:unknown] llus subtilis complete genome (section gene name: ipc-31d; similar to] t] >gp:[GI:e1184599:g2636218] unknown] [OR:Bacillus subtilis] e (section 19 of 21): from 3597091to
ORF Name	NTID	AAID	NT AA score probability
A17503000979_24642963_c2_458	75	3847	216 71 81 0.0019
sp:[LN:COPP_HELFE] [AC:O32620] [GN:COPROTEIN (COPPER ION BINDING PROTEIN)] >gp:[GI:e353967:g2660543] [LN:HFAJ19361] binding protein] [OR:Helicobacter fecopA, copP genes and two ORF's.] [SPORF Name AI7503000979_24647558_c1_352] [SP:03 32] [AC: lis] [DB	2620] AJ001 3:genp	[DB:swissprot] 932] [GN:copP] [FN:divalent cation ept-bct1] [DE:Helicobacter felis ftsH,
Description			JLJ L
NO-HIT			
ORF Name A17503000979_24665932_c3_496	NTID	AAID 3849	NT AA LengthLength score probability 840 279 477 2.1e-45
	>pir:[LN og ywkE] 320D] [A omosomal :P45873] 019] [AC	I:S554 [GN:] C:Z49 DNA [LE:	38] [AC:S55438:D70061] ywkE] [OR:Bacillus subtilis] 782] [GN:ywkE] [OR:Bacillus subtilis] (region 320-321 degrees).] [NT:product 24695] [RE:25561] [DI:direct] 22:AL009126] [GN:ywkE] [FN:unknown]

ORF Name	NTID	AAID	NT AA LengthLength score probability
AI7503000979_2541301_c3_505	78	3850	351 116 180 6.3e-14
Description			
pir:[LN:G70070] [AC:G70070] [PN:hypsubtilis] [DB:pir2] >gp:[GI:e1184584 [GN:ywzB] [FN:unknown] [OR:Bacillus complete genome (section 19 of 21): [DI:complement] >gp:[GI:e1184584:g26]	1:g263620 subtilis from 359 536203] [[DB:genp	03] [L s] [DB 97091t [LN:BS pept]	N:BSUB0019] [AC:Z99122:AL009126] :genpept-bct1] [DE:Bacillus subtilis o 3809700.] [LE:182127] [RE:182357] UB0019] [AC:Z99122:AL009126] [GN:ywzB] [DE:Bacillus subtilis complete genome
ORF Name	NTID	AAID	NT AA score probability
AI7503000979_25422081_c3_475	79	3851	204 67 84 0.014
Description		<u> </u>	
sp:[LN:ETF1_FOWP1] [AC:P21966] [GN:FTRANSCRIPTION FACTOR 70 KD SUBUNIT] [AC:F35216] [PN:FPD6 protein] [CL:v[OR:fowlpox virus] [DB:pir2] >gp:[GI[FN:Vaccinia D6 homolog] [OR:Fowlpox sequence (Hind III fragment).] [SP:F	[SP:P219 vaccinia [:g61229] v virus])66] [virus [LN: [DB:g	DB:swissprot] >pir:[LN:F35216] early transcription factor 70K chain] POFPHIND] [AC:X17202] [GN:ORF FPD6] enpept-vrl] [DE:Fowlpox virus DNA
ORF Name	NTID	AAID	NT AA score probability
AI7503000979_26173800_c1_387	80	3852	186 61
Description			JL L
NO-HIT			
ORF Name	NTID	AAID	NT AA Score probability
AI7503000979_26182767_c2_426	81	3853	123 40
Description	<u></u>		d
NO-HIT			
ORF Name	NTID	AAID	NT AA score probability
A17503000979_26212756_f2_137	82	3854	1477 2.3e-151
Description			
sp:[LN:YWNE_BACSU] [AC:P71040] [GN:Y			
KD PROTEIN IN SPOIIQ-MTA INTERGENIC	REGION]	[SP:P	71040] [DB:swissprot] >pir:[LN:G70063]
[AC:G70063] [PN:cardiolipin synthas cardiolipin synthetase] [OR:Bacillus	e nomolo	g ywni	i GN:ywnE CL:Bacillus probable
[LN:BSUB0019] [AC:Z99122:AL009126] [
[DB:genpept-bct1] [DE:Bacillus subti	lis comp	lete	genome (section 19 of 21): from
3597091to 3809700.] [NT:similar to c	ardiolip	in sy	nthase] [SP:P71040] [LE:164628]
[RE:166076] [DI:direct] >gp:[GI:e269	549:g159	2701]	[LN:BSUEROP] [AC:Y08559] [PN:Unknown]
[GN:ywnE] [OR:Bacillus subtilis] [DB			
<pre>downstream DNA.] [NT:Product similar [LE:5155] [RE:6603] [DI:complement]</pre>			
[AC:Z99122:AL009126] [GN:ywnE] [FN:u			
[DE:Bacillus subtilis complete genom			

[NT:similar to cardiolipin synthase] [SP:P71040] [LE:164628] [RE:166076] [DI:direct]

ODE Name	NIMITO		NT AA		
ORF Name	NTID	AAID	LengthLength score probability		
A17503000979_26360260_c3_520	83	3855	375 124 416 6.1e-39		
Description gp:[GI:e1340260:g3850850] [LN:SAU164 [OR:Staphylococcus aureus] [DB:genpe partial kdpC gene and 4ORF's.] [LE:2	pt-bct1]	[DE:	Staphylococcus aureus dpj, alr genes,		
ORF Name	NTID	AAID	NT AA probability		
AI7503000979_26751542_f1_73	84	3856	492 163 577 5.3e-56		
hypothetical protein HI0491] [OR:Hel [LN:AE000532] [AC:AE000532:AE000511]	icobacte [PN:cor enpept-h	er pyl nserve oct2]	d hypothetical protein] [GN:HP0105] [DE:Helicobacter pylori 26695 section		
ORF Name AI7503000979_26757677_c2_408 Description	NTID 85	<u>AAID</u> 3857	NT AA LengthLength score probability 1377 458 2163 4.6e-224		
gp:[GI:e1352473:g3892895] [LN:SAARGFEMD] [AC:Y15477] [PN:phosphoglucosamine-mutase] [GN:glmM] [OR:Staphylococcus aureus] [DB:genpept-bct1] [DE:Staphylococcus aureus argI, glmM genes and ORF1 and ORF2.] [LE:2968] [RE:4323] [DI:direct] >gp:[GI:e283110:g1684749] [LN:SAFEMD] [AC:Y09570] [GN:femD] [OR:Staphylococcus aureus] [DB:genpept-bct1] [DE:S.aureus femD gene.] [LE:511] [RE:1866] [DI:direct]					
ORF Name	NTID	AAID	NT AA LengthLength score probability		
AI7503000979_2775462_c1_394	86	3858	522 173 331 6.2e-30		
Description gp:[GI:e1340259:g3850849] [LN:SAU16431] [AC:Y16431] [PN:hypothetical protein] [GN:ORF4] [OR:Staphylococcus aureus] [DB:genpept-bct1] [DE:Staphylococcus aureus dpj, alr genes, partial kdpC gene and 4ORF's.] [LE:2477] [RE:2971] [DI:direct]					
ORF Name	NTID	AAID	NT AA LengthLength score probability		
AI7503000979_2925275_f1_20	87	3859	141 46		
Description NO-HIT					
ORF Name AI7503000979_29307312_c3_470 Description	NTID 88	<u>AAID</u> 3860	NT AA LengthLength 165 54		

NO-HIT

ORF Name NTID A	AID <u>NT AA</u> LengthLength score probability
AI7503000979_29695252_c3_497 89 38	361 1098 365 665 2.5e-65
Description Sp:[LN:YWLC_BACSU] [AC:P39153] [GN:YWLC:IPC-29] [DE:HYPOTHETICAL 37.0 KD PROTEIN IN SPOIIR-GLY [DB:swissprot] >pir:[LN:I40476] [AC:I40476:H70] [protein ywlC:SUA5 homolog ipc-29d] [GN:ywlC:ipn [protein ywlC:SUA5 homolog ipc-29d] [DI:direct] >pn [protein ywlC:IA0476] [DI:direct] >pn [protein ywlC:IA0476	CC INTERGENIC REGION] [SP:P39153] O061:S49358] [PN:conserved hypothetical oc-29d] [CL:Bacillus subtilis conserved dis] [DB:pir2] >gp:[GI:g556881] Charomyces cerevisiae SUA5] [GN:ipc-29d] B. subtilis spoII-R, glyC and upp genes.] GP:[GI:e1184601:g2636220] [LN:BSUB0019] OR:Bacillus subtilis] [DB:genpept-bct1] A 19 of 21): from 3597091to 3809700.] [SP:P39153] [LE:194934] [RE:195974] U:BSUB0019] [AC:Z99122:AL009126] [GN:ywlC] Ot] [DE:Bacillus subtilis complete genome contact and complete genome contact and contact and complete genome contact and
	AID NT AA Score probability BEET 189 62
	>pir:[LN:D31482] [AC:D31482] ::H+-transporting ATP synthase chain I] [2] >gp:[GI:g142557] [LN:BACATPA] [ase b subunit] [OR:Bacillus megaterium] [0, pCAH1.3, and pWPC208] [e i,a,c,b,delta,alpha,gamma,beta]
	LengthLength score probability 164 891 296 1229 4.3e-125

gp:[GI:e1352471:g3892893] [LN:SAARGFEMD] [AC:Y15477] [PN:hypothetical protein]
[OR:Staphylococcus aureus] [DB:genpept-bct1] [DE:Staphylococcus aureus argI, glmM genes and ORF1 and ORF2.] [NT:orf1] [LE:1198] [RE:2007] [DI:direct]

ORF Name	NTID	AAID	<u>NT</u> <u>AA</u> LengthLeng	CCOYA	probability
AI7503000979_33412800_c2_429	93	3865	882 293	135	2e-08
Description pir: [LN:B69595] [AC:B69595] [PN:special [PN:B69595] [PN:special [PN:Bacillus subtilis] [DB:pir2] > gp [AC:D84432:D82370] [PN:BltD] [OR:Baccillus [PN:BltD] [DR:Baccillus [PN:BltD] [DB [PN:BltD] [DB [PN:B1] [LE] > gp: [GI:e1183889:g2635105] [LN:BSUBOcacetyltransferase] [GN:bltD] [OR:Baccillus subtilis complete genome [NT:alternate gene name: bmr2D, bmtD]	:[GI:d10 illus su :genpept :811] [F 014] [A0 illus su e (secti	013033 ubtilis c-bct1 RE:126 C:2991 ubtilis	g1303698] [SR:Baci [DE:Baci]] [DI:comp .7:AL009126] [DB:genr of 21): fr	[LN:BACJI illus subti ilus subti olement] o] [PN:spe pept-bct1]	H642] Cilis Cilis DNA, 283 Kb Ermine/spermidine [EC:2.3.1] Sito 2812870.]
ORF Name	NTID	AAID	<u>NT</u> <u>AA</u> LengthLeng	CCOYA	probability
A17503000979_3361326_c2_403	94	3866	<u> 207 68</u>		
Description NO-HIT					
ORF Name	NTID	AAID	<u>NT AA</u> LengthLeng		probability
AI7503000979_33673776_c2_435	95	3867	297 98	312 6	.4e-28
gp:[GI:g4193373] [LN:AF072894] [AC:A: [OR:Listeria monocytogenes] [DB:genpetranscription terminator Rho(rho) genglycosylationprotein GtcA (gtcA) and [LE:848] [RE:1093] [DI:direct]	ept-bct2 ne, part	l [DE ial co	Listeria m ls; and wal	nonocytoge .l teichoi	enes 4b1 putative .c acid
ORF Name	NTID	AAID	<u>NT AA</u> LengthLeng	SCOTE	probability
A17503000979_33751260_c1_372	96	3868	471 156	448 2	.5e-42
Description pir: [LN:D70065] [AC:D70065] [PN:(3R] dehydratase, ywpB] [GN:ywpB] [CL:(3R] dehydratase] [OR:Bacillus subtilis] [LN:BSUB0019] [AC:Z99122:AL009126] [GR:genpept-bct1] [DE:Bacillus subtilis] 3597091to 3809700.] [NT:similar to hy [RE:146122] [DI:complement] >gp:[GI:GR:ywpB] [OR:Bacillus subtilis] [DB ywp[B,C,D,E,F,G,H,I,J] and ywqAgenes protein)] [LE:3199] [RE:3597] [DI:dir [AC:Z99122:AL009126] [GN:ywpB] [FN:ur [DE:Bacillus subtilis complete genome [NT:similar to hydroxymyristoyl-(acy: [DI:complement]	R)-hydro [EC: 4.2 GN:ywpB] lis comp ydroxymy e289141: genpept .] [NT:s rect] >g nknown] e (secti	pxymyr: [FN:: [FN:: [olete gyristoy g1763] [-bct1] similar gp:[GI: [OR:Ba	stoyl-[acy [DB:pir2] nknown] [Cenome (sec 1-(acyl ca 03] [LN:BS [DE:B.sub to hydrox e1184543:g cillus sub of 21): fr	yl carrier ygp:[GI:e PR:Bacillu tion 19 c rrier pro Z83337] [tilis mbl rymyristoy [2636162] tilis] [D Tom 359709	r protein] el184543:g2636162] as subtilis] of 21): from otein)] [LE:145724] [AC:Z83337] a, flh[O,P], rapD, el-(acyl carrier [LN:BSUB0019] oB:genpept] olto 3809700.]
ORF Name	NTID	AAID	<u>NT AA</u> LengthLeng	SCOTE	probability
AT7503000979_33870312_c3_522	97	3869	792 263		.1e-123
Description					
<pre>gp:[GI:e279935:g1934991] [LN:SASIGFAG [OR:Staphylococcus aureus] [DB:genpey [RE:3444] [DI:direct] >gp:[GI:e28500G [GN:sigB] [OR:Staphylococcus aureus] sigB genes.] [LE:2687] [RE:3457] [DI:</pre>	pt-bct1] D:g17297 [DB:gen	[DE:S 97] [I pept-k	.aureus si N:SAUSIGB]	gB gene.] [AC:Y099	[LE:2674] 29] [PN:sigma-B]

ORF Name	NTID	AAID	NT AA LengthLength score probability			
A17503000979_34062928_c3_521	98	3870	1083 360 1321 7.7e-135			
Description						
<pre>gp:[GI:e1340261:g3850851] [LN:SAU16431] [AC:Y16431] [PN:alr protein] [GN:alr] [OR:Staphylococcus aureus] [DB:genpept-bct1] [DE:Staphylococcus aureus dpj, alr genes, partial kdpC gene and 4ORF's.] [LE:3400] [RE:4548] [DI:direct]</pre>						
ORF Name	NTID	AAID	NT AA score probability			
A17503000979_34181277_c2_461	99	3871	501 166 299 1.5e-26			
Description	T					
	pt-bct1] [DE:	1] [PN:hypothetical protein] [GN:ORF2] Staphylococcus aureus dpj, alr genes, [DI:direct]			
ORF Name	NTID	AAID	NT AA score probability			
A17503000979_34187702_c2_407	100	3872	936 311 1173 3.7e-119			
Description						
gp:[GI:e1352472:g3892894] [LN:SAARGFEMD] [AC:Y15477] [PN:hypothetical protein] [GN:orf2] [OR:Staphylococcus aureus] [DB:genpept-bct1] [DE:Staphylococcus aureus argI, glmM genes and ORF1 and ORF2.] [LE:2009] [RE:2941] [DI:direct]						
ORF Name	NTID	AAID	NT AA score probability			
AI7503000979_34197275_£2_176	101	3873	126 41			
Description						
NO-HIT						
ORF Name	NTID	AAID	NT AA LengthLength score probability			
A17503000979_34250327_c1_396	102	3874	363 120 540 4.5e-52			
Description						
gp:[GI:e279931:g1934987] [LN:SASIGFACB] [AC:Y07645] [GN:ORF1] [OR:Staphylococcus aureus] [DB:genpept-bct1] [DE:S.aureus sigB gene.] [LE:72] [RE:434] [DI:direct]						
ORF Name	NTID	AAID	NT AA score probability			
A17503000979_34589010_c1_348	103	3875	246 81			
Description						
NO-HIT						

NT ORF Name NTID score probability LengthLength AI7503000979 34611067 ±3 302 104 540 Description sp:[LN:YWJG BACSU] [AC:P06629] [GN:YWJG] [OR:BACILLUS SUBTILIS] [DE:HYPOTHETICAL 19.1 KD PROTEIN IN SPOOF-PYRG INTERGENIC REGION (ORFS)] [SP:P06629] [DB:swissprot] >pir:[LN:140471] [AC:140471:S55424:E70060] [PN:hypothetical protein ywjG:spoOF protein] [GN:ywjG:spoOF] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:g853763] [LN:BSDNA320D] [AC:Z49782] [GN:ywjG] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:B.subtilis chromosomal DNA (region 320-321 degrees).] [SP:P06629] [LE:11169] [RE:11690] [DI:complement] >gp:[GI:g40177] [LN:BSSP00] [AC:V00105:J01549] [GN:spoOF] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis gene required at an early stage of sporulation.(gene code spoOF).] [SP:P06629] [LE:541] [RE:1062] [DI:direct] >gp:[GI:e1184620:g2636239] [LN:BSUB0019] [AC:Z99122:AL009126] [GN:ywjG] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 19 of 21): from 3597091to 3809700.] [SP:P06629] [LE:212053] [RE:212574] [DI:direct] >gp:[GI:e1186215:g2636251] [LN:BSUB0020] [AC:Z99123:AL009126] [GN:ywjG] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 20 of 21): from 3798401to 4010550.] [SP:P06629] [LE:10743] [RE:11264] [DI:direct] >gp:[GI:e1184620:g2636239] [LN:BSUB0019] [AC:Z99122:AL009126] [GN:ywjG] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept] [DE:Bacillus subtilis complete genome (section 19 of 21): from 3597091to 3809700.] [SP:P06629] [LE:212053] [RE:212574] [DI:direct] NTAΑ Length Length score ORF Name NTID probabi<u>lity</u> AAID AI7503000979 34615700 f2 193 105 819 2.8e-08 Description gp:[GI:g1293846] [LN:CELC42D8] [AC:U56966] [GN:C42D8.3] [OR:Caenorhabditis elegans] [SR:Caenorhabditis elegans strain=Bristol N2] [DB:genpept-inv1] [DE:Caenorhabditis elegans cosmid C42D8.] [NT:coded for by C. elegans cDNA yk30b3.5; coded for by] [LE:8907:9377:9844:10820] [RE:9056:9564:10206:10976] [DI:directJoin] NT AA ORF Name NTID AAID score probability LengthLength AI7503000979_34646926_£2_206 106 3878 732 243 585 7.6e-57 Description sp:[LN:DEOD_ACTPL] [AC:P94164] [GN:DEOD] [OR:ACTINOBACILLUS PLEUROPNEUMONIAE] [SR:, HAEMOPHILUS PLEUROPNEUMONIAE] [EC:2.4.2.1] [DE:(PNP)] [SP:P94164] [DB:swissprot] >gp:[GI:g1732037] [LN:APU55016] [AC:U55016] [PN:purine nucleoside phosphorylase] [GN:deoD] [FN:cleavage of guanosine or inosine to respective] [OR:Actinobacillus pleuropneumoniae] [DB:genpept-bct1] [EC:2.4.2.1] [DE:Actinobacillus pleuropneumoniae heat-shock 10 protein GroES (mopB), heat-shock 60 protein GroEL (mopA), purine nucleoside phosphorylase (deoD) genes, complete cds, alcohol dehydrogenase (adhE)

gene, partial cds.] [LE:176] [RE:898] [DI:direct]

NTAΑ Length Length score ORF Name NTID AAID probability AI7503000979_35354656_f1_108 107 3879 228 75 Description NO-HIT

$rac{ ext{NTID}}{ ext{NAME}}$ $rac{ ext{NTID}}{ ext{LengthLength}}$ $rac{ ext{Score}}{ ext{prob}}$	pability
17503000979_35647783_£1_103 108 3880 690 229 181 4.9e-1	.4
Description pir:[LN:F71082] [AC:F71082] [PN:hypothetical protein PH0924] [GN:PH0924] [OR:Pyrococcus horikoshii] [DB:pir2] >gp:[GI:d1030963:g3257337] [LN:AP0000 [AC:AP000004:AB009494:AB009495:AB009496:AB009497:AB009498:AB009499] [PN:12 hypothetical protein] [GN:PH0924] [OR:Pyrococcus horikoshii] [SR:Pyrococcu (strain:OT3) DNA] [DB:genpept-bct1] [DE:Pyrococcus horikoshii OT3 genomic 777001-994000 nt. position(4/7).] [LE:55768] [RE:56154] [DI:direct]	004] 28aa long 1s horikoshii
ORF Name NTID AAID NT AA Score prok	pability
Description	
sp:[LN:ATPG_BACME] [AC:P20602] [GN:ATPG] [OR:BACILLUS MEGATERIUM] [EC:3.6.5] SYNTHASE GAMMA CHAIN,] [SP:P20602] [DB:swissprot] >pir:[LN:G31482] [AC:G314] [PN:H+-transporting ATP synthase, gamma chain] [CL:H+-transporting ATP synthain] [OR:Bacillus megaterium] [EC:3.6.1.34] [DB:pir2] >gp:[GI:g142560] [AC:M20255:J04455:M18352:M23924] [PN:ATP synthase gamma subunit] [OR:Bacilmegaterium] [SR:B.megaterium (QM B1551) DNA, clones pWSB100, pCAH1.3, and [DB:genpept-bct1] [DE:B.megaterium ATP synthase i,a,c,b,delta,alpha,gamma,andepsilon subunit genes, complete cds, and ORF.] [LE:4467] [RE:5324] [DI:	1482] ithase gamma [LN:BACATPA] llus pWPC208] beta
ORF Name NTID AAID NT AA score prob	pability
A17503000979_36225052_c3_502	3
Description pir:[LN:I39786] [AC:I39786] [PN:H+-transporting ATP synthase, C chain] [O pir:[LN:I39786] [AC:I39786] [PN:H+-transporting ATP synthase, C chain] [O [CL:H+-transporting ATP synthase lipid-binding protein] [OR:Bacillus firmu [EC:3.6.1.34] [DB:pir2] >gp:[GI:g142570] [LN:BACATPSYNB] [AC:M84713] [PN:F subunit] [GN:atpE] [FN:proton translocation] [OR:Bacillus firmus] [SR:Baci DNA] [DB:genpept-bct1] [DE:Bacillus firmus ATP synthase a and c subunit gently and complete cds.] [NT:putative] [LE:299] [RE:511] [DI:direct]	ns] ATP synthase c .llus firmus
ORF Name NTID AAID LengthLength score prob	ability
Length Length Score Prob	ability
A17503000979_36363432_f1_85	ability
A17503000979_36363432_f1_85	ability
MID HAID LengthLength Score Probability LengthLength Length Lengt	ability
NTID AAID LengthLength Score Probe	

gp:[GI:g5410434] [LN:AF134170] [AC:AF134170] [PN:gag] [OR:Dictyostelium discoideum]
[DB:genpept] [DE:Dictyostelium discoideum retrotransposon TRE3-B, complete sequence.] [LE:139] [RE:1347] [DI:direct]

ORF Name	NTID	AAID	NT AA LengthLength
A17503000979_4063202_c3_499	113	3885	1239 412 1563 1.8e-160
Description			
[OR:Bacillus subtilis] [DB:genpept-b genome (section 19 of 21): from 3597 ipc-34d] [SP:P39148] [LE:191155] [RE [LN:BSUB0019] [AC:Z99122:AL009126] [FN:glycine/serine/threonine metabol [EC:2.1.2.1] [DE:Bacillus subtilis c	[DB:sw ne hydr lyC] [[DB:pir transfe II-R, g 96:g263 [GN:gl: ct1] [E 091to 3 :192402 PN:serii ism] [O omplete	issprooxymet! CL:gly 2] >gp rase] lyC and 6215] yA] [F] C:2.1.: 809700] [DI: ne hyd: genome	ch] >pir:[LN:I40483] chyltransferase, glyA:serine ccine hydroxymethyltransferase] c:[GI:g556886] [LN:BSSPORUPP] [GN:glyC] [OR:Bacillus subtilis] cd upp genes.] [SP:P39148] [LE:5499] [LN:BSUB0019] [AC:Z99122:AL009126] [N:glycine/serine/threonine metabolism] 2.1] [DE:Bacillus subtilis complete c.] [NT:alternate gene name: glyC, complement] >gp:[GI:e1184596:g2636215] droxymethyltransferase] [GN:glyA]
ORF Name A17503000979_40712_f2_210	NTID	<u>AAID</u>	NT AA score probability LengthLength 71
Description NO-HIT			
ORF Name	NTID	AAID	NT AA LengthLength score probability
AI7503000979_4072135_c3_490	115	3887	1467 488 870 4.8e-87
Description		•	
	pir2] > l prote Mycobac	gp:[GI in Rv0: terium	:e304956:g1871596] [LN:MTCY8D5] 223c] [GN:Rv0223c] [OR:Mycobacterium tuberculosis H37Rv complete genome;
ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000979_4079511_f1_3	116	3888	147 48
Description			
NO-HIT			
ORF Name	NTID	AAID	NT AA LengthLength score probability
AI7503000979_409556_f1_111	117	3889	135 44 112 2.5e-06
	>gp:[G: 466:AB0 R:Pyroce [DE:Pyro	I:d1030 09467: <i>I</i> occus l	0234:g3256608] [LN:AP000001] AB009468:AB009469] [PN:235aa long horikoshii] [SR:Pyrococcus horikoshii s horikoshii OT3 genomic DNA, 1-287000

ORF Name	NTID	AAID	NT AA score probability			
A17503000979_4178218_c1_353	118	3890	675 224 81 0.0066			
Description						
gp:[GI:g1131502] [LN:PBU42580] [AC:U	ept-vrl]	[DE:	Paramecium bursaria Chlorella virus 1,			
ORF Name	NTID	AAID	NT AA LengthLength score probability			
A17503000979_4346926_c2_450	119	3891	123 40			
Description						
NO-HIT						
ORF Name	NTID	AAID	NT AA Length Length score probability			
A17503000979_4728558_c2_409	120	3892	1860 619 2088 4.1e-216			
Description						
pir:[LN:B69633] [AC:B69633] [PN:L-glutamine-D-fructose-6-phosphate amidotransferase glmS] [GN:glmS] [CL:glutaminefructose-6-phosphate aminotransferase (isomerizing)] [GR:Bacillus subtilis] [DB:pir2] >gp:[GI:d1034037:g3599596] [LN:AB006424] [AC:AB006424] [PN:L-GLUTAMINE-D-FRUCTOSE-6-PHOSPHATE AMIDO] [GN:gcaA] [OR:Bacillus subtilis] [SR:Bacillus subtilis (strain:168) DNA] [DB:genpept-bct1] [DE:Bacillus subtilis genomic DNA, 70 kb region between 17 and 23degree.] [LE:3246] [RE:5048] [DI:direct] >gp:[GI:g726480] [LN:BSU21932] [AC:U21932:D21198] [PN:L-glutamine-D-fructose-6-phosphate] [GN:gcaA] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis L-glutamine-D-fructose-6-phosphateamidotransferase (gcaA) gene, complete cds.] [LE:312] [RE:2114] [DI:direct] >gp:[GI:e1182111:g2632445] [LN:BSUB0001] [AC:Z99104:AL009126] [PN:L-glutamine-D-fructose-6-phosphate] [GN:glmS] [OR:Bacillus subtilis] [DB:genpept-bct1] [EC:2.6.1.16] [DE:Bacillus subtilis complete genome (section 1 of 21): from 1 to213080.] [NT:alternate gene name: gcaA, ybxD] [SP:P39754] [LE:200263] [RE:202065] [DI:direct] >gp:[GI:e1182129:g2632463] [LN:BSUB0002] [AC:Z99105:AL009126] [PN:L-glutamine-D-fructose-6-phosphate] [GN:glmS] [OR:Bacillus subtilis] [DB:genpept-bct1] [EC:2.6.1.16] [DE:Bacillus subtilis complete genome (section 2 of 21): from 194651 to415810.] [NT:alternate gene name: gcaA, ybxD] [SP:P39754] [LE:5613] [RE:7415] [DI:direct]						
ORF Name	NTID	AAID	NT AA score probability			
A17503000979_476567_c1_355	121	3893	567 188 292 8.5e-26			
subtilis] [EC:2.7.7.6] [DB:pir1] >gp subtilis] [SR:B.subtilis (strain 168) [DE:B.subtilis RNA polymerase delta s protein (ttg start codon)] [LE:101] [LN:BSDNA320D] [AC:Z49782] [PN:RNA polymerase] subtilis] [DB:genpept-bct1] [DE:B.subtilis] [DB:genpept-bct1] [DE:B.subtilis] [SP:P12464] [LE >gp:[GI:e1186217:g2636253] [LN:BSUB00]	A SUBUNI 55422:H6 -directe :[GI:g14) DNA, coubunit [RE:622] olymeras otilis co :8717] [020] [AC	T,] [9 9698] d RNA 3456] llone m (rpoE) [DI:c e delt chromos RE:923 :Z9912 DB:ger	SP:P12464] [DB:swissprot]] [PN:DNA-directed RNA polymerase, polymerase delta chain] [OR:Bacillus [LN:BACRPOE] [AC:M21677] [OR:Bacillus mML11] [DB:genpept-bct1]) gene, complete cds.] [NT:rpoE direct] >gp:[GI:g853761] ta subunit] [GN:rpoE] [OR:Bacillus somal DNA (region 320-321 degrees).] 38] [DI:direct] 23:AL009126] [PN:RNA polymerase (delta npept-bct1] [EC:2.7.7.6] [DE:Bacillus			
[LE:13195] [RE:13716] [DI:complement]						

ORF Name	NTID	AAID	Length Length score	probability		
AI7503000979_4869213_c3_519	122	3894		3.4e-128		
Description		L	JJ			
<pre>gp:[GI:e1340258:g3850848] [LN:SAU164 [OR:Staphylococcus aureus] [DB:genpe partial kdpC gene and 40RF's.] [LE:9</pre>	ept-bct1] [DE:	Staphylococcus aureu			
ORF Name	NTID	AAID	NT AA LengthLength	probability		
AI7503000979_4881262_f3_293	123	3895	171 56			
Description						
NO-HIT						
ORF Name	NTID	AAID	NT AA LengthLength	probability		
A17503000979_4881302_f2_191	124	3896	150 49			
Description NO-HIT						
ORF Name	NTID	AAID	NT AA LengthLength	probability		
AI7503000979_4901712_c1_370	125	3897	1428 475 1999 1	.1e-206		
[DE:(VEG31)] [SP:P37809] [DB:swisspr [PN:H+-transporting ATP synthase, be ATP synthase alpha chain: H+-transporting ATP synthase alpha chain: H+-transporting ATP synthase subtilis] [OR:Bacillus subtilis] [EC:3.6.1.34] [AC:Z28592] [PN:ATP synthase subunity and subtilis (16, b, delta, alpha, gamma, beta, epsilops, beta, alpha, gamma, beta, epsilops, beta) [GN:atpD] [OR:Bacillus subtilis subtilis complete genome (section 19) [LE:183456] [RE:184877] [DI:compleme [AC:Z99122:AL009126] [PN:ATP synthase [DB:genpept] [EC:3.6.1.34] [DE:Bacil from 3597091to 3809700.] [SP:P37809]	eta chair prting A' [DB:pir beta] [SB) atpas [On.] [SI [O19] [AC [is] [DB [of 21) [ont] >gp [se (subur [lus subt	n (atp IP syn r2] >g [GN:at se gen P:P378 C:Z991 :genpe :from :[GI:e nit be tilis	D) [GN:atpD] [CL: thase alpha chain ho p:[GI:g433991] [LN:E pD] [OR:Bacillus sub es for ATP synthase 09] [LE:5520] [RE:69 22:AL009126] [PN:ATP pt-bct1] [EC:3.6.1.3 3597091to 3809700.] 1184587:g2636206] [L ta)] [GN:atpD] [OR:B complete genome (sec	H+-transporting omology] BSATPASE] otilis] subunits i, a, c 041] [DI:direct] O synthase (subunit 04] [DE:Bacillus [SP:P37809] W:BSUB0019] Bacillus subtilis] Stion 19 of 21):		
ORF Name	NTID	AAID	NT AA LengthLength score	probability		
A17503000979_4962802_c2_421	126	3898	1305 434 1591 1	.9e-163		
Description gp:[GI:g4336793] [LN:AF105341] [AC:AF105341] [PN:pyrimidine nucleoside phosphorylase] [GN:pdp1] [OR:Listeria monocytogenes] [DB:genpept-bct2] [DE:Listeria monocytogenes threonine dehydratase (thd1) gene, partialcds; alpha acetolactate decarboxylase gene, complete cds; andpyrimidine nucleoside phosphorylase (pdp1) gene, partial cds.] [LE:1749] [RE:>3010] [DI:complement]						
ORF Name	NTID	AAID	NT AA LengthLength	probability		
AI7503000979_5078177_c1_365	127	3899	1062 353 1169 9	.9e-119		
Description gp:[GI:g1773355] [LN:SAU81973] [AC:U aureus] [DB:genpept-bct2] [DE:Staphy Cap5Pgenes, complete cds.] [NT:putat [RE:16944] [DI:direct]	lococcus	s aure	us capsule gene clus	ter Cap5A through		

ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000979_5111502_c2_454	128	3900	924 307 417 4.8e-39
Description		L	
homolog yqjG] [GN:yqjG] [CL:stage I protein homology] [OR:Bacillus subti [LN:BACJH642] [AC:D84432:D82370] [PN (strain:JH642(trpC2 PheA1)) DNA] [DE region containing skin element.] [LE >gp:[GI:e1185657:g2634823] [LN:BSUB0	G69963] III sport Lis] [Di S:YqjG] S:genpept S:234919 D13] [Acctl] [Di [NT:sim:	[AC:Gulation B:pir2 [OR:Bat-bct1] [RE:C:Z991 E:Baciilar to	G69963] [PN:lipoprotein SpoIIIJ-like on protein:stage III sporulation 2] >gp:[GI:d1013293:g1303958] acillus subtilis] [SR:Bacillus subtilis 1] [DE:Bacillus subtilis DNA, 283 Kb:235746] [DI:complement] 116:AL009126] [GN:yqjG] [FN:unknown] illus subtilis complete genome (section
ORF Name A17503000979_5131927_f1_93	NTID	<u>AAID</u>	<u>LengthLength</u>
Description			
NO-HIT			
ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000979_5318785_c1_395	130	3902	[156] [51]
Description NO-HIT			
ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000979_5319213_c2_442	131	3903	417 138
SYNTHASE EPSILON CHAIN,] [SP:P37812] [AC:I40369:G69591:S39257] [PN:H+-tr [GN:atpC] [CL:H+-transporting ATP s [EC:3.6.1.34] [DB:pir2] >gp:[GI:g433 subunit epsilon] [GN:atpC] [OR:Bacil (168) atpase genes for ATP synthase epsilon.] [SP:P37812] [LE:6965] [RE:[LN:BSUB0019] [AC:Z99122:AL009126] [OR:Bacillus subtilis] [DB:genpept-b genome (section 19 of 21): from 3597 [DI:complement] >gp:[GI:e1184586:g26]	[DB:swinter ansported property of the content of th	issprofing ATI epsilo N:BSATI tilis] s i, a DI:dire synthas C:3.6.1 809700 [LN:BSI acillus ion 19	TP synthase, epsilon chain (atpC)] lon chain] [OR:Bacillus subtilis] TPASE] [AC:Z28592] [PN:ATP synthase] [DB:genpept-bct1] [DE:B.subtilis] a, c, b, delta, alpha, gamma, beta, rect] >gp:[GI:el184586:g2636205] ase (subunit epsilon)] [GN:atpC] .1.34] [DE:Bacillus subtilis complete D.] [SP:P37812] [LE:183034] [RE:183432] GUB0019] [AC:Z99122:AL009126] [PN:ATP us subtilis] [DB:genpept] [EC:3.6.1.34] e of 21): from 3597091to 3809700.]
ORF Name	NTID	AAID	NT AA score probability
A17503000979_582760_c2_463	132	3904	<u>LengthLength</u> 150 9.5e-11
Description		<u> </u>	
gp:[GI:e1340262:g3850852] [LN:SAU164	pt-bct1]	[DE:S	[PN:hypothetical protein] [GN:ORF7] Staphylococcus aureus dpj, alr genes, [BI:direct]

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability	
AI7503000979_5895301_c1_378	133	3905	948	315	603	9.4e-59	7
Description		<u> </u>					_
sp:[LN:THID_HAEIN] [AC:P44697] [GN:T [EC:2.7.4.7] [DE:(HMP-P KINASE)] [SP] [PN:hypothetical protein HI0416] [[OR:Haemophilus influenzae] [DB:pir2 [PN:phosphomethylpyrimidine kinase ([DB:genpept-bct2] [DE:Haemophilus in genome.] [NT:similar to GB:AE000511 [DI:direct]	:P44697 CL:phos] >gp:[(thiD)] fluenza] [DB: phomet GI:g15 [GN:HI e Rd s	swisspr hylpyri 73390] 0416] { ection	cot] >p midine [LN:U3 [OR:Hae 40 of	ir:[LN:phosp 2725] mophil 163 of	::I64151] [AC:I64 hate kinase] [AC:U32725:L4202 us influenzae Rd the complete	:3]
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability	
AI7503000979_6442192_f2_181	134	3906		43			
Description							
NO-HIT							
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability	
A17503000979_6454635_c2_431	135	3907	912	303	1173	3.7e-119	7
Description							_
sp:[LN:ALF1_BACSU] [AC:P13243] [GN:F. [EC:4.1.2.13] [DE:PROBABLE FRUCTOSE-IDB:swissprot] >pir:[LN:D32354] [AC:PN:fructose-bisphosphate aldolase, orfy-tsr:fructose-1,6-bisphosphate aldolase II] [OR:Bacillus subtilis] [LN:BACSPO0FA] [AC:M22039] [PN:fructIOR:Bacillus subtilis] [SR:Bacillus [DB:genpept-bct1] [EC:4.1.2.13] [DE:andfructose-bisphosphate aldolase (of [DI:direct] >gp:[GI:g853765] [LN:BSDIO aldolase] [GN:tsr] [OR:Bacillus subtIDNA (region 320-321 degrees).] [SP:P>gp:[GI:e1184618:g2636237] [LN:BSUBO [PN:fructose-1,6-bisphosphate aldolase subtilis] [DB:genpept-bct1] [EC:4.1.1. (section 19 of 21): from 3597091to 3 [SP:P13243] [LE:210475] [RE:211332] [LN:BSUBO020] [AC:Z99123:AL009126] [FN:glycolysis] [OR:Bacillus subtilis subtilis complete genome (section 20 gene name: tsr, fba] [SP:P13243] [LE>gp:[GI:e1184618:g2636237] [LN:BSUBO020] [PN:fructose-1,6-bisphosphate aldolase subtilis] [DB:genpept] [EC:4.1.2.13] of 21): from 3597091to 3809700.] [NT [LE:210475] [RE:211332] [DI:complement the subtilis] [DB:genpept] [EC:4.1.2.13] of 21): from 3597091to 3809700.] [NT [LE:210475] [RE:211332] [DI:complement the subtilis] [RE:211332]	BISPHOSI S55426:I fbaA:30I ldolase: [EC:4.1 ose-bisp subtilis Bacillus rfY-tsr) NA320D ilis] [I 13243] 019] [AG se] [GN 2.13] [I 809700.] [DI:comp PN:fruct s] [DB:G of 21) :9165] 019] [AG se] [GN: cof 21) :100:10000000000000000000000000000000	PHATE D32354 K phosp C [GN:: 2.13] phosphose (strices subted) generated [AC:Zental] [LE:12ental] [LE:12ental] [Entered [ALDOLAS :E32354 phoprot fbaA] [DB:pi ate ald ains JH illis s s, comp 49782] pept-bc 411] [R 22:AL00 [FN:gl illus s alterna t] >gp: ,6-bisp t-bct1] 379840 022] [D 022:AL00 [FN:gl subtil	E 1,] ::D4183 ein [CL:fr r1] >g lolase] 642 an pooF, lete c [PN:fr tt] [D E:1326 9126] ycolys ubtili te gen [GI:el hospha [EC:4 1to 40 I:comp 9126] ycolys is com	[SP:P1 5:B696 uctose p:[GI: [GN:c d UOTO CTP sy ds.] [uctose E:B.su 8] [DI is] [C s comp e name 186213 te ald .1.2.1 10550. lement is] [C plete	3243] 21] -bisphosphate g460911] rfY-tsr] 550) DNA] nthetase (ctrA), LE:3270] [RE:412 biphosphate btilis chromosom :direct] R:Bacillus lete genome : tsr, fba] :g2636249] olase] [GN:fbaA] 3] [DE:Bacillus] [NT:alternate] R:Bacillus genome (section in	al
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability	
A17503000979_6645393_c1_342	136	3908		54			
Description			- I				

NO-HIT

ORF Name	NTID	AAID LengthLength score probability
A17503000979_6681316_f2_197	137	3909 135 44
Description		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
AI7503000979_6721877_f2_194	138	3910 153 50
Description	•	
NO-HIT	···-	
ORF Name	NTID	AAID NT AA score probability
AI7503000979_6906576_c1_371	139	3911 1296 431 1429 2.8e-146
Description		
		rlglucosamine 1-carboxyvinyltransferase
murA] [GN:murA] [CL:UDP-N-acetylglu [OR:Bacillus subtilis] [DB:pir2] sqn		e 1-carboxyvinyltransferase MurZ] [6830:g1648861] [LN:BSATPC] [AC:Z81356]
[PN:UDP-N-acetylglucosamine] [GN:mur		
		[DI:direct] >gp:[GI:e1184582:g2636201]
[LN:BSUB0019] [AC:Z99122:AL009126] [J-acetylglucosamine] [GN:murA] s subtilis] [DB:genpept-bct1] [EC:2.5.1.7]
-		on 19 of 21): from 3597091to 3809700.]
[LE:179914] [RE:181224] [DI:compleme	nt] >gp:	[GI:e1184582:g2636201] [LN:BSUB0019]
[AC:Z99122:AL009126] [PN:UDP-N-acety		
		pept] [EC:2.5.1.7] [DE:Bacillus subtilis 97091to 3809700.] [LE:179914] [RE:181224]
		(Ma.101221)
[DI:complement]		
[DI:Complement]	· · · · · · · · · · · · · · · · · · ·	NT AA
ORF Name	NTID	AAID NT AA score probability
	NTID	AATD — gcore probability
ORF Name		Length Length score probability
ORF Name AI7503000979_7240675_c2_460 Description sp:[LN:MURF_BACSU] [AC:P96613] [GN:M [DE:(D-ALANYL-D-ALANINE-ADDING ENZYM [AC:F69662] [PN:UDP-N-acetylmuramoy	URF] [OR E)] [SP:	AAID LengthLength score probability 3912 1374 457 976 2.8e-98 2:BACILLUS SUBTILIS] [EC:6.3.2.15] P96613] [DB:swissprot] >pir:[LN:F69662] D-glutamyl-2, 6-diaminopimelate-D-al murF]
ORF Name A17503000979_7240675_c2_460 Description sp:[LN:MURF_BACSU] [AC:P96613] [GN:M [DE:(D-ALANYL-D-ALANINE-ADDING ENZYM [AC:F69662] [PN:UDP-N-acetylmuramoy [GN:murF] [OR:Bacillus subtilis] [D	URF] [OR E)] [SP: lalanyl- B:pir2]	LengthLength Score probability 3912 1374 457 976 2.8e-98 E:BACILLUS SUBTILIS] [EC:6.3.2.15] P96613] [DB:swissprot] >pir:[LN:F69662] D-glutamyl-2, 6-diaminopimelate-D-al murF] >gp:[GI:d1020047:g1881267] [LN:AB001488]
ORF Name A17503000979_7240675_c2_460 Description sp:[LN:MURF_BACSU] [AC:P96613] [GN:M [DE:(D-ALANYL-D-ALANINE-ADDING ENZYM [AC:F69662] [PN:UDP-N-acetylmuramoy [GN:murF] [OR:Bacillus subtilis] [D [AC:AB001488] [GN:ydbQ] [OR:Bacillus	URF] [OR E)] [SP: lalanyl- B:pir2] subtili	LengthLength Score probability 3912 1374 457 976 2.8e-98 E:BACILLUS SUBTILIS] [EC:6.3.2.15] P96613] [DB:swissprot] >pir:[LN:F69662] D-glutamyl-2, 6-diaminopimelate-D-al murF] >gp:[GI:d1020047:g1881267] [LN:AB001488] s] [SR:Bacillus subtilis (strain:168) DNA]
ORF Name AI7503000979_7240675_c2_460 Description sp:[LN:MURF_BACSU] [AC:P96613] [GN:M [DE:(D-ALANYL-D-ALANINE-ADDING ENZYM [AC:F69662] [PN:UDP-N-acetylmuramoy [GN:murF] [OR:Bacillus subtilis] [D [AC:AB001488] [GN:ydbQ] [OR:Bacillus [DB:genpept-bct1] [DE:Bacillus subtiregionbetween 35 and 47 degree.] [NT	URF] [OR E)] [SP: lalanyl- B:pir2] subtili lis geno	LengthLength Score Probability
ORF Name AI7503000979_7240675_c2_460 Description sp: [LN:MURF_BACSU] [AC:P96613] [GN:M [DE: (D-ALANYL-D-ALANINE-ADDING ENZYM [AC:F69662] [PN:UDP-N-acetylmuramoy [GN:murF] [OR:Bacillus subtilis] [D [AC:AB001488] [GN:ydbQ] [OR:Bacillus [DB:genpept-bct1] [DE:Bacillus subtiregionbetween 35 and 47 degree.] [NT [LE:42447] [RE:43820] [DI:direct] >g	URF] [OR E)] [SP: lalanyl- B:pir2] subtili lis geno :PROBABL p:[GI:e1	LengthLength Score Probability
ORF Name AI7503000979_7240675_c2_460 Description sp:[LN:MURF_BACSU] [AC:P96613] [GN:M [DE:(D-ALANYL-D-ALANINE-ADDING ENZYM [AC:F69662] [PN:UDP-N-acetylmuramoy [GN:murF] [OR:Bacillus subtilis] [D [AC:AB001488] [GN:ydbQ] [OR:Bacillus [DB:genpept-bct1] [DE:Bacillus subtiregionbetween 35 and 47 degree.] [NT [LE:42447] [RE:43820] [DI:direct] >g [AC:Z99106:AL009126] [PN:UDP-N-acety	URF] [OR E)] [SP: lalanyl- B:pir2] subtili lis geno :PROBABL p:[GI:e1 lmuramoy	LengthLength Score Probability
ORF Name AI7503000979_7240675_c2_460 Description sp: [LN:MURF_BACSU] [AC:P96613] [GN:M [DE: (D-ALANYL-D-ALANINE-ADDING ENZYM [AC:F69662] [PN:UDP-N-acetylmuramoy [GN:murF] [OR:Bacillus subtilis] [D [AC:AB001488] [GN:ydbQ] [OR:Bacillus [DB:genpept-bct1] [DE:Bacillus subtiregionbetween 35 and 47 degree.] [NT [LE:42447] [RE:43820] [DI:direct] >g [AC:Z99106:AL009126] [PN:UDP-N-acety [FN:peptidoglycan biosynthesis] [OR: [DE:Bacillus subtilis complete genome	URF] [OR E)] [SP: lalanyl- B:pir2] subtili lis geno :PROBABL p:[GI:e1] lmuramoy Bacillus e (secti	LengthLength Score Probability
ORF Name AI7503000979_7240675_c2_460 Description sp:[LN:MURF_BACSU] [AC:P96613] [GN:M [DE:(D-ALANYL-D-ALANINE-ADDING ENZYM [AC:F69662] [PN:UDP-N-acetylmuramoy [GN:murF] [OR:Bacillus subtilis] [D [AC:AB001488] [GN:ydbQ] [OR:Bacillus [DB:genpept-bct1] [DE:Bacillus subtiregionbetween 35 and 47 degree.] [NT [LE:42447] [RE:43820] [DI:direct] >g [AC:Z99106:AL009126] [PN:UDP-N-acety [FN:peptidoglycan biosynthesis] [OR:	URF] [OR E)] [SP: lalanyl- B:pir2] subtili lis geno :PROBABL p:[GI:e1] lmuramoy Bacillus e (secti	LengthLength Score Probability
ORF Name AI7503000979_7240675_c2_460 Description sp: [LN:MURF_BACSU] [AC:P96613] [GN:M [DE: (D-ALANYL-D-ALANINE-ADDING ENZYM [AC:F69662] [PN:UDP-N-acetylmuramoy [GN:murF] [OR:Bacillus subtilis] [D [AC:AB001488] [GN:ydbQ] [OR:Bacillus [DB:genpept-bct1] [DE:Bacillus subtiregionbetween 35 and 47 degree.] [NT [LE:42447] [RE:43820] [DI:direct] >g [AC:Z99106:AL009126] [PN:UDP-N-acety [FN:peptidoglycan biosynthesis] [OR: [DE:Bacillus subtilis complete genome	URF] [OR E)] [SP: lalanyl- B:pir2] subtili lis geno :PROBABL p:[GI:e1] lmuramoy Bacillus e (secti	LengthLength Score Probability
ORF Name AI7503000979_7240675_c2_460 Description sp:[LN:MURF_BACSU] [AC:P96613] [GN:M [DE:(D-ALANYL-D-ALANINE-ADDING ENZYM [AC:F69662] [PN:UDP-N-acetylmuramoy [GN:murF] [OR:Bacillus subtilis] [D [AC:AB001488] [GN:ydbQ] [OR:Bacillus [DB:genpept-bct1] [DE:Bacillus subti regionbetween 35 and 47 degree.] [NT [LE:42447] [RE:43820] [DI:direct] >g [AC:Z99106:AL009126] [PN:UDP-N-acety [FN:peptidoglycan biosynthesis] [OR: [DE:Bacillus subtilis complete genom [NT:alternate gene name: ydbQ] [SP:P	URF] [OR E)] [SP: lalanyl- B:pir2] subtili lis geno :PROBABL p:[GI:e1 lmuramoy Bacillus e (secti	LengthLength Score Probability
ORF Name AI7503000979_7240675_c2_460 Description sp:[LN:MURF_BACSU] [AC:P96613] [GN:M [DE:(D-ALANYL-D-ALANINE-ADDING ENZYM [AC:F69662] [PN:UDP-N-acetylmuramoy [GN:murF] [OR:Bacillus subtilis] [D [AC:AB001488] [GN:ydbQ] [OR:Bacillus [DB:genpept-bct1] [DE:Bacillus subtiregionbetween 35 and 47 degree.] [NT [LE:42447] [RE:43820] [DI:direct] >g [AC:Z99106:AL009126] [PN:UDP-N-acety [FN:peptidoglycan biosynthesis] [OR: [DE:Bacillus subtilis complete genom [NT:alternate gene name: ydbQ] [SP:PORF Name	URF] [OR E)] [SP: lalanyl- B:pir2] subtili lis geno :PROBABL p:[GI:e1] lmuramoy Bacillus e (secti 96613] [LengthLength Score Probability
ORF Name AI7503000979_7240675_c2_460 Description sp: [LN:MURF_BACSU] [AC:P96613] [GN:M [DE: (D-ALANYL-D-ALANINE-ADDING ENZYM [AC:F69662] [PN:UDP-N-acetylmuramoy [GN:murF] [OR:Bacillus subtilis] [D [AC:AB001488] [GN:ydbQ] [OR:Bacillus [DB:genpept-bct1] [DE:Bacillus subtiregionbetween 35 and 47 degree.] [NT [LE:42447] [RE:43820] [DI:direct] >g [AC:Z99106:AL009126] [PN:UDP-N-acety [FN:peptidoglycan biosynthesis] [OR: [DE:Bacillus subtilis complete genom [NT:alternate gene name: ydbQ] [SP:P ORF Name AI7503000979_7292200_c1_351	URF] [OR E)] [SP: lalanyl- B:pir2] subtili lis geno :PROBABL p:[GI:e1] lmuramoy Bacillus e (secti 96613] [LengthLength Score Probability
ORF Name AI7503000979_7240675_c2_460 Description sp:[LN:MURF_BACSU] [AC:P96613] [GN:M [DE:(D-ALANYL-D-ALANINE-ADDING ENZYM [AC:F69662] [PN:UDP-N-acetylmuramoy [GN:murF] [OR:Bacillus subtilis] [D [AC:AB001488] [GN:ydbQ] [OR:Bacillus [DB:genpept-bct1] [DE:Bacillus subtiregionbetween 35 and 47 degree.] [NT [LE:42447] [RE:43820] [DI:direct] >g [AC:Z99106:AL009126] [PN:UDP-N-acety [FN:peptidoglycan biosynthesis] [OR:[DE:Bacillus subtilis complete genom [NT:alternate gene name: ydbQ] [SP:PORF Name AI7503000979_7292200_c1_351 Description	URF] [OR E)] [SP: lalanyl- B:pir2] subtili lis geno :PROBABL p:[GI:e1] lmuramoy Bacillus e (secti 96613] [LengthLength Score Probability
ORF Name AI7503000979_7240675_c2_460 Description sp: [LN:MURF_BACSU] [AC:P96613] [GN:M [DE: (D-ALANYL-D-ALANINE-ADDING ENZYM [AC:F69662] [PN:UDP-N-acetylmuramoy [GN:murF] [OR:Bacillus subtilis] [D [AC:AB001488] [GN:ydbQ] [OR:Bacillus [DB:genpept-bct1] [DE:Bacillus subtiregionbetween 35 and 47 degree.] [NT [LE:42447] [RE:43820] [DI:direct] >g [AC:Z99106:AL009126] [PN:UDP-N-acety [FN:peptidoglycan biosynthesis] [OR: [DE:Bacillus subtilis complete genom [NT:alternate gene name: ydbQ] [SP:PORF Name AI7503000979_7292200_c1_351 Description NO-HIT	URF] [OR E)] [SP: lalanyl- B:pir2] subtili lis geno :PROBABL p:[GI:e1 lmuramoy Bacillus e (Secti 96613] [NTID 141	LengthLength LengthLength LengthLength P76 2.8e-98 LEBACILLUS SUBTILIS] [EC:6.3.2.15] P96613] [DB:swissprot] >pir:[LN:F69662] D-glutamyl-2, 6-diaminopimelate-D-al murf] >gp:[GI:d1020047:g1881267] [LN:AB001488] s] [SR:Bacillus subtilis (strain:168) DNA] me sequence, 148 kb sequence of the LE UDP-N-ACETYLMURAMOYLALANYL-D-GLUTAMYL-2,] 182423:g2632757] [LN:BSUB0003] Plalanyl-D-glutamyl-2,6-] [GN:murf] subtilis] [DB:genpept-bct1] [EC:6.3.2.15] on 3 of 21): from 402751 to611850.] LE:106191] [RE:107564] [DI:direct] AAID NT AA LengthLength Score probability 3913 126 41
ORF Name AI7503000979_7240675_c2_460 Description sp: [LN:MURF_BACSU] [AC:P96613] [GN:M [DE: (D-ALANYL-D-ALANINE-ADDING ENZYM [AC:F69662] [PN:UDP-N-acetylmuramoy [GN:murF] [OR:Bacillus subtilis] [D [AC:AB001488] [GN:ydbQ] [OR:Bacillus [DB:genpept-bct1] [DE:Bacillus subtiregionbetween 35 and 47 degree.] [NT [LE:42447] [RE:43820] [DI:direct] >g [AC:Z99106:AL009126] [PN:UDP-N-acety [FN:peptidoglycan biosynthesis] [OR: [DE:Bacillus subtilis complete genom [NT:alternate gene name: ydbQ] [SP:PORF Name AI7503000979_7292200_c1_351 Description NO-HIT ORF Name	URF] [OR E)] [SP: lalanyl- B:pir2] subtili lis geno :PROBABL p:[GI:e1 lmuramoy Bacillus e (Secti 96613] [NTID 141 NTID	LengthLength 976 2.8e-98

ORF Name	NTID	AAID	NT AA LengthLength	probability
A17503000979_9800787_t2_216 Description	143	3915	129 42	
NO-HIT				
			NT AA	
ORF Name	NTID	AAID	NT AA LengthLength score	probability
AI7503000980_10006507_£3_164	144	3916	153 50	
Description				
NO-HIT				
ORF Name	NTID	AAID	NT AA LengthLength	probability
A17503000980_10736002_c1_211	145	3917	1521 506 2313 5	5.9e-240
Description				
>gp:[GI:g567884] [LN:STA5SRR] [AC:L3 aureus] [SR:Staphylococcus aureus (c [DE:Staphylococcus aureus lysyl-tRNA genes, 5S ribosomal RNA (5S rRNA) geRNA (23S rRNA) gene.] [LE:176] [RE:1	lone li sythet ne, 16S	brary: ase ge riboso	ATCC 12600) DNA] [I ene, complete cds,tra emal RNA (16S rRNA) c	DB:genpept-bct1] ansfer RNA (tRNA)
ORF Name	NTID	AAID	NT AA LengthLength score	probability
A17503000980_11210316_c1_203	146	3918	195 64 166 1	9e-12
Description				•
sp:[LN:YABO_BACSU] [AC:P37557] [GN:Y PROTEIN IN MFD-DIVIC INTERGENIC REGI				
[AC:S66089:F69739] [PN:conserved hy				
subtilis] [DB:pir2] >gp:[GI:d1005836	:g46744	8] [LN	:BAC180K] [AC:D26185	[PN:unknown]
[OR:Bacillus subtilis] [SR:Bacillus				
[DB:genpept-bct1] [DE:B. subtilis DN [LE:131477] [RE:131737] [DI:direct]				
[AC:Z99104:AL009126] [GN:yab0] [FN:u				
[DE:Bacillus subtilis complete genom	e (sect	ion 1	of 21): from 1 to213	080.] [NT:similar
to hypothetical proteins] [SP:P37557	[LE:6	7875]	[RE:68135] [DI:direc	et]
ORF Name	NTID	AAID	NT AA LengthLength	probability
A17503000980_1292767_c3_243	147	3919		

ORF Name	NTID AAI	ID NT AA score probability
A17503000980_12929625_c1_186	148 392	20 810 269 819 1.2e-81
Description	L L	
KD PROTEIN IN XPAC-ABRB INTERGENIC R [AC:S66062:A69738] [PN:signal pepti subtilis] [DB:pir2] >gp:[GI:d1005810 [OR:Bacillus subtilis] [SR:Bacillus [DB:genpept-bct1] [DE:B. subtilis DN [LE:105257] [RE:106084] [DI:direct] [AC:Z99104:AL009126] [GN:yaaT] [FN:u	EGION] [SP: dase II hom :g467422] [subtilis (s A, 180 kilo >gp:[GI:e11 nknown] [OR e (section	[LN:BAC180K] [AC:D26185] [PN:unknown] sub_species:Marburg, strain:168) DNA] bbase region of replication origin.] 181965:g2632299] [LN:BSUB0001] R:Bacillus subtilis] [DB:genpept-bct1] 1 of 21): from 1 to213080.] [NT:similar
ORF Name	NTID AAI	ID NT AA score probability
A17503000980_13759688_c1_184	149 392	21 615 204 527 1.1e-50
<pre>subtilis] [DB:pir2] >gp:[GI:d1005806 [OR:Bacillus subtilis] [SR:Bacillus [DB:genpept-bct1] [DE:B. subtilis DN [LE:102759] [RE:103397] [DI:direct] [AC:Z99104:AL009126] [PN:thymidylate</pre>)] [SP:P375 kinase tmk] :g467418] [subtilis (s A, 180 kilo >gp:[GI:e11 kinase] [G acillus sub	[GN:tmk] [CL:dTMP kinase] [OR:Bacillus [LN:BAC180K] [AC:D26185] [PN:unknown] [Sub_species:Marburg, strain:168) DNA] [Dbase region of replication origin.] [B1961:g2632295] [LN:BSUB0001] [GN:tmk] [OR:Bacillus subtilis] [Dtilis complete genome (section 1 of 21):
ORF Name	NTID AAI	- Length Length
A17503000980_14460015_f1_55 Description	150 392	22 141 46
NO-HIT		
ORF Name A17503000980_16484577_f2_119 Description	NTID AAI	Length Length
NO-HIT		
ORF Name AI7503000980_16522641_f2_76	NTID AAI	Length Length
Description	C4] [70 750	0000041 [DW 252cc languages 1
<pre>gp:[GI:d1045277:g5106180] [LN:AP0000 protein] [GN:APE2475] [OR:Aeropyrum [DB:genpept] [DE:Aeropyrum pernix general</pre>	pernix] [SR	R:Aeropyrum pernix (strain:K1) DNA]

OWL:AP00000385 percent identity:66.667] [LE:151512] [RE:152573] [DI:direct]

		NI	AA 7		
ORF Name	NTID	7 7 TD	thLength sco	ore probabi	lity
A17503000980_16594202_c2_237	153	3925 480	159 38:	3 1.9e-35	
Description					
sp:[LN:HPPK_BACSU] [AC:P29252] [GN:F (6-HYDROXYMETHYL-7,8-DIHYDROPTERIN F [DB:swissprot] >pir:[LN:S66109] [AC:[PN:2-amino-4-hydroxy-6-hydroxymethy] pyrophosphokinase,:6-hydroxymethyl-78-dihydro-6-hydroxymethylpterin pyroc[CL:2-amino-4-hydroxy-6-hydroxymethy] pyrophosphokinase:2-amino-4-hydroxy-homology] [OR:Bacillus subtilis] [EC [LN:BAC180K] [AC:D26185] [PN:7,8-dih [GN:folk] [OR:Bacillus subtilis] [SR strain:168) DNA] [DB:genpept-bct1] [replication origin.] [LE:150545] [RE [LN:BSUB0001] [AC:Z99104:AL009126] [FN:dihydrofolate biosynthesis] [OR:[DE:Bacillus subtilis complete genom [RE:87446] [DI:direct]	PYROPHOSE S66109: It I dihydro I, 8-dihydro Phosphol Vldihydro 6-hydro 2:2.7.6.3 1ydro-6-l 2:Bacillu [DE:B. su E:151048] [PN:7,8-d Bacillus	HOKINASE) 37854:F696 pteridine ropterin p inase] [Gh pteridine ymethyldih] [DB:pir2 ydroxymeth s subtilis btilis DNA [DI:direc ihydro-6-h subtilis]	(PPPK)] [SP 626] pyrophosphok N:folK] nydropteridi 2] >gp:[GI:d nylpterin-py s (sub_speci A, 180 kilob ct] >gp:[GI: nydroxymethy [DB:genpep	ne pyrophosp 1005856:g467 rophosphokin es:Marburg, ase region of e1182012:g26 lpterin] [GN t-bct1] [EC:	7468] n] of 532346] V:folK] :2.7.6.3]
ORF Name AI7503000980 190875 f1 6	NTID	AAID Leng	thLength sco	re probabi	lity
	154	3926 210	69 70	0.028	
Description pir:[LN:S69873] [AC:S69873] [PN:hyp cerevisiae] [DB:pir2] [MP:13L]	othetica	l protein	YML009w-a]	[OR:Sacchard	omyces
ORF Name	NTID	AAID NT	AA thLength	re probabi	lity
AI7503000980_194142_c1_194	155	3927 138			
Description					.
pir:[LN:S66080] [AC:S66080:I40018:C6 pyrophosphorylase gcaD:cell division [CL:N-acetylglucosamine-1-phosphate >gp:[GI:d1005827:g467439] [LN:BAC180 division] [GN:tms26] [OR:Bacillus su strain:168) DNA] [DB:genpept-bct1] [replication origin.] [LE:119952] [RE [LN:BSUB0001] [AC:Z99104:AL009126] [[GN:gcaD] [FN:peptidoglycan and lipo [DB:genpept-bct1] [EC:2.7.7.23] [DE: 21): from 1 to213080.] [NT:alternate [RE:57720] [DI:direct]	protein uridyltr [AC:E btilis] [DE:B. su [:121322] PN:UDP-N polysaco Bacillus	tms26:tms ansferase] 26185] [PN [SR:Bacill btilis DNA [DI:direc -acetylglu haride] [C subtilis	s protein] [6] [OR:Bacille N:temperature Lus subtilis A, 180 kilobact] >gp:[GI:6] acosamine py: OR:Bacillus acomplete gen	GN:gcaD:tms2 us subtilis] e sensitive (sub_specie ase region o e1181983:g26 rophosphoryl subtilis] nome (sectio	[DB:pir2] cell es:Marburg, of [32317] ase]
ORF Name	NTID		th Length Sco	re probabi	lity
AI7503000980_20335927_f2_69	156	3928 153	50		

ORF Name	NTID	AAID	NT AA score	probability
A17503000980 211687 c3 261		3929	LengthLength	
Description	157	3929	897 298 985 3.1	Le-99
sp:[LN:YACC_BACSU] [AC:P37565] [GN: KD PROTEIN IN FTSH-CYSK INTERGENIC [AC:S66101:F69740] [PN:conserved h hypothetical protein sll1988] [OR:B >gp:[GI:d1005848:g467460] [LN:BAC18 [SR:Bacillus subtilis (sub_species:subtilis DNA, 180 kilobase region o [DI:direct] >gp:[GI:e1182004:g26323 [FN:unknown] [OR:Bacillus subtilis] genome (section 1 of 21): from 1 to [SP:P37565] [LE:79877] [RE:80752] [REGION] ypotheti acillus 0K] [AC: Marburg, f replic 38] [LN: [DB:gen 213080.]	[SP:P3 cal pr subtil D26185 strai ation BSUB00 pept-b [NT:s	7565] [DB:swissprot] : otein yacC] [GN:yacC] is] [DB:pir2]] [PN:unknown] [OR:Bac n:168) DNA] [DB:genper origin.] [LE:143479] 01] [AC:Z99104:AL00912 ct1] [DE:Bacillus subt	>pir:[LN:S66101]] [CL:conserved cillus subtilis] pt-bct1] [DE:B. [RE:144354] 26] [GN:yacC] tilis complete
ORF Name	NTID	AAID	NT AA LengthLength score	probability
AI7503000980_21579131_f3_157	158	3930	123 40	
Description				
NO-HIT				
ORF Name A17503000980_22694002_c3_258 Description sp:[LN:HPRT_BACSU] [AC:P37472] [GN:: [DE:(HGPRTASE)] [SP:P37472] [DB:swi. [PN:hypoxanthine phosphoribosyltran phosphoribosyltransferase hprT] [GN [OR:Bacillus subtilis] [EC:2.4.2.8] [AC:D26185] [PN:hypoxanthine-guanin subtilis] [SR:Bacillus subtilis (su [DB:genpept-bct1] [DE:B. subtilis Di [LE:139944] [RE:140486] [DI:direct] [AC:Z99104:AL009126] [PN:hypoxanthin [FN:purine salvage] [OR:Bacillus su subtilis complete genome (section 1 [RE:76884] [DI:direct]	ssprot] sferase, :hprT] [DB:pir e phosph b_specie NA, 180 >gp:[GI ne-guani btilis] of 21):	>pir:[hprT: [CL:hy 2] >gp oribos s:Marb kiloba :e1182 ne pho [DB:ge	BACILLUS SUBTILIS] [ECLN:S66098] [AC:S66098: hypoxanthine-guanine poxanthine phosphoribot: [GI:d1005845:g467457] yltransferase] [GN:hprurg, strain:168) DNA] se region of replication of replication of sphoribosyltransferase sphoribosyltransferase npept-bct1] [EC:2.4.2.1 to213080.] [SP:P3747	e:E69642] psyltransferase] [LN:BAC180K] pt] [OR:Bacillus don origin.] 30001] pt] [GN:hprT] 8] [DE:Bacillus 72] [LE:76342]
ORF Name	NTID	AAID	NT AA LengthLength score p	robability
A17503000980_23445130_c1_207	160	3932	417 138 134 4.1	e-08
Description				
<pre>gp:[GI:e1182000:g2632334] [LN:BSUB00 [OR:Bacillus subtilis] [DB:genpept-] 1 of 21): from 1 to213080.] [NT:sim:</pre>	bct1] [D	E:Baci	llus subtilis complete	genome (section

[DI:direct]

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000980_23601702_c1_190	161	3933	891	296	919	3.1e-92
Description						
sp:[LN:KSGA_BACSU] [AC:P37468] [GN:K [DE:DIMETHYLTRANSFERASE)] [SP:P37468] [PN:dimethyladenosine transferase [GN:ksgA] [CL:rRNA (adenine-N6-)-me >gp:[GI:d1005819:g467431] [LN:BAC180] resistance] [GN:ksgA] [OR:Bacillus s (sub_species:Marburg, strain:168) DN kilobase region of replication origit >gp:[GI:e1181975:g2632309] [LN:BSUB0 transferase] [GN:ksgA] [FN:high leve [DB:genpept-bct1] [EC:2.1.1] [DE:Bfrom 1 to213080.] [SP:P37468] [LE:50] [DB:s ksgA:hi thyltra K] [AC: ubtilis A] [DB: n.] [LE 001] [A l kasug acillus	wissprogh levensferas D26185] [SR:Figenpept ::114240 C:Z9910 amycin subtil	ot] >pi el kasg se] [OR [PN:h Bacillu -bct1] D] [RE: D4:AL00 resist Lis com	er:[LN: gamycin gamyci	s66071 resis lus su evel ka :ilis s. subt s] [DI: [PN:di [OR:Ba genome	cl] [AC:S66071:A69649 stance protein ksgA] stallis [DB:pir2] ssgamycin cliis DNA, 180 direct] methyladenosine scillus subtilis]
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000980_23605438_c2_238	162	3934		47		
Description			-,		•	
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000980_23631327_c3_250	163	3935	966	321	1225	1.1e-124
gp:[GI:g2289093] [LN:CAU76387] [AC:U [OR:Corynebacterium ammoniagenes] [Di ammoniagenes N-acetyl glucoseamine 1 cds, and PRPP-synthetase(prs) gene,	B : genpe -phosph	pt-bct1 ateurio	l] [EC: dyltran [LE:3	2.7.6. sferas 21] [R	1] [DE se (glm	::Corynebacterium NU) gene, partial
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000980_23642135_c1_185	164	3936	357	118	389	4.5e-36
Description sp:[LN:YAAQ_BACSU] [AC:P37538] [GN:YAKD PROTEIN IN XPAC-ABRB INTERGENIC RITECT [AC:S66059:G69737] [PN:conserved hypothetical proteins] [SR:Bacillus subtilis] [DB:pir2] >gp:[GI:d1005807] [OR:Bacillus subtilis] [SR:Bacillus [DB:genpept-bct1] [DE:B. subtilis DN:[LE:103471] [RE:103800] [DI:direct] [AC:Z99104:AL009126] [GN:yaaQ] [FN:waward [DE:Bacillus subtilis complete genome to hypothetical proteins] [SP:P37538]	EGION] potheti :g46741 subtili A, 180 >gp:[GI nknown] e (sect	[SP:P37] cal pro 9] [LN: s (sub_ kilobas :e11819 [OR:Ba ion 1 c	otein y BAC180 specie se regi 962:g26 acillus of 21):	DB:swi aaQ] [K] [AC s:Marb on of 32296] subti from	ssprot GN:yaa ::D2618 ourg, s replic [LN:B lis] [1 to21] >pir:[LN:S66059] Q] [OR:Bacillus 5] [PN:unknown] train:168) DNA] ation origin.] SUB0001] DB:genpept-bct1] 3080.] [NT:similar
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000980_23860952_c1_206	165	3937		293		4.5e-27
Description					_	
gp:[GI:g4981097] [LN:AE001733] [AC:Alprotein] [GN:TM0579] [OR:Thermotogal section 45 of 136 of the complete get 59.32;] [LE:230] [RE:1474] [DI:direction 45 of 136 of the complete get 59.32;]	maritim nome.]	a] [DB:	genpep	t-bct2] [DE:	Thermotoga maritima

ORF Name	NTID	AAID	NT LengthI	<u>AA</u> Length	score	probability
A17503000980_24350953_±3_120	166	3938	132	43	72	0.017
Description pir:[LN:G71244] [AC:G71244] [PN:hype [OR:Pyrococcus horikoshii] [DB:pir2] [AC:AP000001:AB009465:AB009464:AB009469] hypothetical protein] [GN:PH0217] [OR:CREATE CONTROL OF CON	>gp:[G] 466:AB00 R:Pyroco [DE:Pyro	:d103 9467:2 occus l	0229:g32 AB009468 horikosh s horiko	256603 B:AB00 nii] [oshii] [LN: 9469] SR:Pyr	AP000001] [PN:106aa long Tococcus horikoshii
ORF Name	NTID	AAID	NT LengthL	AA ength	score	probability
A17503000980_24647936_c3_256	167	3939	408	135	172	4.4e-13
Description gp:[GI:g4090866] [LN:AF023181] [AC:Almonocytogenes] [DB:genpept-bct2] [DE coupling factor (mfdL),low temperatus homolog(divL) genes, complete cds.]	:Listeri re requi	a mond	ocytoger t B prot	nes tr	anscri ltrB),	ption-repair
ORF Name	NTID	AAID	<u>NT</u> LengthL	AA ength	score	probability
A17503000980_24649092_c2_236 Description	168	3940		125	351	4.7e-32
sp:[LN:FOLB_STAHA] [AC:Q59920] [GN:FO [EC:4.1.2.25] [DE:DIHYDRONEOPTERIN AN [DB:swissprot] >gp:[GI:g1118003] [LN aldolase] [GN:folQ] [OR:Staphylococcu [DE:Staphylococcus haemolyticus cyste (folQ) genes, partial cds, anddihydro [NT:DHNA] [LE:1467] [RE:>1724] [DI:di	LDOLASE, :SHU4076 us haemo eine syn opteroat	(DHN) [A0] Slyticu Sthase	A) (FRAG C:U40768 1s] [DB: A (cysk	GMENT) B] [PN genpe C) and] [SP: :dihyd pt-bct dihydr	Q59920] roneopterin 2] [EC:4.1.2.25] oneopterin aldolase
ORF Name	NTID	AAID	<u>NT</u> LengthL	<u>AA</u> ength	score	probability
A17503000980_24663892_c3_251	169	3941	1881	526	1224	1.5e-124
Description sp:[LN:MFD_BACSU] [AC:P37474] [GN:MFI COUPLING FACTOR (TRCF)] [SP:P37474] [PN:transcription-repair coupling fac coupling protein:DEAD/H box helicase >gp:[GI:d1005832:g467444] [LN:BAC1801 factor] [GN:mfd] [OR:Bacillus subtili strain:168) DNA] [DB:genpept-bct1] [I replication origin.] [LE:124030] [RE: [LN:BSUB0001] [AC:Z99104:AL009126] [I [FN:probably involved in homologous I [DE:Bacillus subtilis complete genome [LE:60428] [RE:63961] [DI:direct]	[DB:swisetor mfd homolog [AC:DE:B. suite 127563] PN:trans DNA] [OR	ssprot] [] [GN: [y] [OF [26185] Bacil] [btilis [DI: [cripti	>pir:[:mfd] [R:Bacill [PN:tr Lus subt s DNA, 1 Lirect] ion-repa	LN:S60 [CL:transcr: lus sulfanscr: lilis .80 ki .>gp:[(lir con	6085] anscri btilis iption (sub_s lobase GI:e11 upling	[AC:S66085:F69657] ption-repair] [DB:pir2] -repair coupling pecies:Marburg, region of 81988:g2632322] factor] [GN:mfd] genpept-bct1]
ORF Name	NTID		NT LengthL	ength-	score	probability
A17503000980_24734661_f1_13 Description	170	3942	138 4	5		
NO-HIT						

ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000980_24790916_c1_208	171	3943	2106 701 2133 6.9e-221
Description sp:[LN:FTSH_BACSU] [AC:P37476] [GN:FDIVISION PROTEIN FTSH HOMOLOG,] [SP:[AC:E69627:S66099] [PN:cell-division heat shock protein ftsH] [GN:ftsH] FtsH/SEC18/CDC48-type ATP-binding do >gp:[GI:d1005846:g467458] [LN:BAC180 [OR:Bacillus subtilis] [SR:Bacillus [DB:genpept-bct1] [DE:B. subtilis DN [LE:140584] [RE:142497] [DI:direct] [AC:Z99104:AL009126] [PN:cell-divisite [FN:involved in major cellular proce [DB:genpept-bct1] [EC:3.4.24] [DE:21): from 1 to213080.] [SP:P37476] [P37476] n prote [CL:cel main ho K] [AC: subtili A, 180 >gp:[GI on prote sses su Bacillu	[DB:swin / ge in / ge l divis mology] D26185] s (sub kilobas :e11820 ein and ch as] s subti	wissprot] >pir:[LN:E69627] eneral stress protein ftsH:class III sion protein ftsH: [OR:Bacillus subtilis] [DB:pir2] [PN:cell division protein] [GN:ftsH] species:Marburg, strain:168) DNA] se region of replication origin.] 002:g2632336] [LN:BSUB0001] d general stress protein] [GN:ftsH] [OR:Bacillus subtilis] tilis complete genome (section 1 of
ORF Name	NTID	AAID	NT AA LengthLength
A17503000980_25662965_c1_195	172	3944	729 242 463 6.4e-44
Description sp:[LN:CTC_BACSU] [AC:P14194] [GN:CT PROTEIN CTC] [SP:P14194] [DB:swisspr		BACILLU	JS SUBTILIS] [DE:GENERAL STRESS
ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000980_26210061_£2_99	173	3945	144 47
Description NO-HIT			
ORF Name	NTID	AAID	NT AA LengthLength
A17503000980_26839660_c3_246	174	3946	855 284 842 4.4e-84
Description gp:[GI:e1386912:g4454322] [LN:SAU132 [OR:Staphylococcus aureus] [DB:genpe (partial).] [NT:ORF2] [LE:1402] [RE:	pt-bct1] [DE:S	Staphylococcus aureus ORF1 and ORF2
ORF Name	NTID	AAID	NT AA LengthLength
A17503000980_2928502_c3_262	175	3947	960 319 1166 2.1e-118
Description sp:[LN:CYSK_BACSU] [AC:P37887] [GN:C [DE:PROTEIN 11) (SOI11)] [SP:P37887]			
ORF Name	NTID	AAID	NT AA LengthLength
A17503000980_29298162_£2_118	176	3948	123 40
Description NO-HIT			
ORF Name	NTID	AAID	NT AA LengthLength
AI7503000980_29307187_c2_213	177	3949	165 54
Description NO-HIT			

ORF Name	NTID	AAID	NT LengthLe	AA ength score	probability
AI7503000980 31446881 c2 215	178	3950			2.1e-15
Description			ــا لـــــــا لــ	الحصا لحصا	
sp:[LN:YABA_BACSU] [AC:P37542] [GN:	ר ממגע [ר	ם. פארד	יווים פווסי	PTITEL [DE.	UVDORUBRICAL 14 1
KD PROTEIN IN XPAC-ABRB INTERGENIC					
[AC:S66063:B69738] [PN:hypothetical					
[DB:pir2] >gp:[GI:d1005811:g467423]					
subtilis] [SR:Bacillus subtilis (sub					
[DB:genpept-bct1] [DE:B. subtilis DI			-		
[LE:106099] [RE:106458] [DI:direct]					
[AC:Z99104:AL009126] [GN:yabA] [FN:	unknown]	[OR:B	acillus s	subtilis] [DB:genpept-bct1]
[DE:Bacillus subtilis complete genor	me (sect	ion 1	of 21): f	From 1 to21	3080.] [SP:P37542]
[LE:42497] [RE:42856] [DI:direct]					
			NT	AA	
ORF Name	NTID	AAID	LengthLe	- 60000	probability
AI7503000980_32615811_f3_166	179	3951	189 62	2	
Description					
NO-HIT			•		
			NTT	77	
ORF Name	NTID	AAID	<u>NT</u> LengthLe	AA ength score	probability
ORF Name A17503000980_33225017_c2_222	NTID	<u>AAID</u>	NT LengthLe	ength score	probability 8.8e-79
			LengthLe	ength score	
A17503000980_33225017_c2_222	180	3952	LengthLe	ength score	8.8e-79
A17503000980_33225017_c2_222 Description	180 PURR] [O	3952 R:BACI	LengthLe	Ength Score 792	8.8e-79 PUR OPERON
Description sp:[LN:PURR_BACSU] [AC:P37551] [GN:REPRESSOR] [SP:P37551] [DB:swissprote [PN:transcription repressor of puris	PURR] [O	3952 R:BACII [LN:S60	LENGTHLE [867 28 LLUS SUBT 5076] [AC [GN:pur	792 DE: C:S66076:D6 CR] [OR:Ba	PUR OPERON 9685] cillus subtilis]
Description sp:[LN:PURR_BACSU] [AC:P37551] [GN:REPRESSOR] [SP:P37551] [DB:swissprot [PN:transcription repressor of purin [DB:pir2] >gp:[GI:d1005824:g467436]	PURR] [O. t] >pir: ne opero: [LN:BAC	3952 R:BACI [LN:S66 n purk]	Length Le	792	PUR OPERON 9685] cillus subtilis] nown] [OR:Bacillus
Description sp:[LN:PURR_BACSU] [AC:P37551] [GN:REPRESSOR] [SP:P37551] [DB:swissprote [PN:transcription repressor of puring [DB:pir2] >gp:[GI:d1005824:g467436] subtilis] [SR:Bacillus subtilis (subtilis)]	PURR] [O t] >pir: ne opero: [LN:BAC o_specie	R:BACII [LN:S60 n purr] 180K] s:Marb	LengthLe B67 28 LLUS SUBT 6076] [AC [AC:D2618 larg, stra	792 [DE: C:S66076:D6 R] [OR:Ba ain:168) DN	PUR OPERON 9685] cillus subtilis] nown] [OR:Bacillus
Description sp:[LN:PURR_BACSU] [AC:P37551] [GN:REPRESSOR] [SP:P37551] [DB:swissprot [PN:transcription repressor of purin [DB:pir2] >gp:[GI:d1005824:g467436] subtilis] [SR:Bacillus subtilis (subtilis [DB:genpept-bct1] [DE:B. subtilis DB	PURR] [O c] >pir: ne opero: [LN:BAC o_specie	R:BACII [LN:S60 n purk] 180K] s:Marbo	LengthLe [867] [28 LLUS SUBT 5076] [AC: [AC:D2618 lurg, stra se region	FILIS] [DE: C:S66076:D6 FR] [OR:Ba B5] [PN:unk ain:168) DN n of replic	PUR OPERON 9685] cillus subtilis] nown] [OR:Bacillus A] ation origin.]
Description sp:[LN:PURR_BACSU] [AC:P37551] [GN:REPRESSOR] [SP:P37551] [DB:swissprotemonth [PN:transcription repressor of puring [DB:pir2] >gp:[GI:d1005824:g467436] subtilis] [SR:Bacillus subtilis (subtilis) [SR:Bacillus subtilis DB:genpept-bct1] [DE:B. subtilis DB [LE:118041] [RE:118898] [DI:direct]	PURR] [O: c] >pir: ne opero: [LN:BAC o_specie NA, 180: >gp:[GI	R:BACII [LN:S66 n purR] 180K] s:Marbi kilobas	LLUS SUBT 5076] [AC:D2618 LC:D2618 Lrg, strase region 980:g2632	FILIS] [DE: C:S66076:D6 CR] [OR:Ba B5] [PN:unk: Ain:168) DN n of replice 2314] [LN:Ba	PUR OPERON 9685] cillus subtilis] nown] [OR:Bacillus A] ation origin.] SUB0001]
Description sp:[LN:PURR_BACSU] [AC:P37551] [GN:IREPRESSOR] [SP:P37551] [DB:swissprote [PN:transcription repressor of puring [DB:pir2] >gp:[GI:d1005824:g467436] subtilis] [SR:Bacillus subtilis (subtilis) [SR:Bacillus subtilis DB:genpept-bct1] [DE:B. subtilis DB [LE:118041] [RE:118898] [DI:direct] [AC:Z99104:AL009126] [PN:transcripts]	PURR] [O]	R:BACII [LN:S60 n purR 180K] s:Marbo kilobas :e11819	LengthLe [867 28 LLUS SUBT 5076] [AC: [AC:D2618 lrg, stra se region 980:g2632 r] [GN:pu	TILIS] [DE: C:S66076:D6 CR] [OR:Ba B5] [PN:unk Bin:168) DN D of replication of re	PUR OPERON 9685] cillus subtilis] nown] [OR:Bacillus A] ation origin.] SUB0001] gative regulation
Description sp:[LN:PURR_BACSU] [AC:P37551] [GN:IREPRESSOR] [SP:P37551] [DB:swissprote [PN:transcription repressor of puring [DB:pir2] >gp:[GI:d1005824:g467436] subtilis] [SR:Bacillus subtilis (subtilis) [SR:Bacillus subtilis DB:genpept-bct1] [DE:B. subtilis DB:[LE:118041] [RE:118898] [DI:direct] [AC:Z99104:AL009126] [PN:transcript:of the purine operon] [OR:Bacillus subtilis of the purine operon]	PURR] [O. t] >pir: ne opero: [LN:BAC o_specie NA, 180 >gp:[GI ional resoubtilis	R:BACII [LN:S60 n purk] 180K] s:Marbo kilobas :e11819 gulato:	LengthLe [867] [28 LLUS SUBT 5076] [AC: [AC:D2618 1rg, stra se region 980:g2632 r] [GN:pu	TILIS] [DE: C:S66076:D6 CR] [OR:Ba B5] [PN:unk ain:168) DN n of replication 2314] [LN:Ba arr] [FN:ne- pott] [DE:Ba	PUR OPERON 9685] cillus subtilis] nown] [OR:Bacillus A] ation origin.] SUB0001] gative regulation acillus subtilis
Description sp:[LN:PURR_BACSU] [AC:P37551] [GN:REPRESSOR] [SP:P37551] [DB:swissprote [PN:transcription repressor of puring [DB:pir2] >gp:[GI:d1005824:g467436] subtilis] [SR:Bacillus subtilis (subtilis) [SR:Bacillus subtilis DB:genpept-bct1] [DE:B. subtilis DB:[LE:118041] [RE:118898] [DI:direct] [AC:Z99104:AL009126] [PN:transcripts of the purine operon] [OR:Bacillus scomplete genome (section 1 of 21):	PURR] [O. t] >pir: ne opero: [LN:BAC o_specie NA, 180 >gp:[GI ional resubtilis from 1 to	R:BACII [LN:S60 n purk] 180K] s:Marbo kilobas :e11819 gulato:] [DB:0	LengthLe [867] [28 LLUS SUBT 5076] [AC: [AC:D2618 1rg, stra se region 980:g2632 r] [GN:pu	TILIS] [DE: C:S66076:D6 CR] [OR:Ba B5] [PN:unk ain:168) DN n of replication 2314] [LN:Ba arr] [FN:ne- pott] [DE:Ba	PUR OPERON 9685] cillus subtilis] nown] [OR:Bacillus A] ation origin.] SUB0001] gative regulation acillus subtilis
Description sp:[LN:PURR_BACSU] [AC:P37551] [GN:IREPRESSOR] [SP:P37551] [DB:swissprote [PN:transcription repressor of puring [DB:pir2] >gp:[GI:d1005824:g467436] subtilis] [SR:Bacillus subtilis (subtilis) [SR:Bacillus subtilis DB:genpept-bct1] [DE:B. subtilis DB:[LE:118041] [RE:118898] [DI:direct] [AC:Z99104:AL009126] [PN:transcript:of the purine operon] [OR:Bacillus subtilis of the purine operon]	PURR] [O. t] >pir: ne opero: [LN:BAC o_specie NA, 180 >gp:[GI ional resubtilis from 1 to	R:BACII [LN:S60 n purk] 180K] s:Marbo kilobas :e11819 gulato:] [DB:0	LengthLe [867] [28 LLUS SUBT 5076] [AC: [AC:D2618 1rg, stra se region 980:g2632 r] [GN:pu	TILIS] [DE: C:S66076:D6 CR] [OR:Ba B5] [PN:unk ain:168) DN n of replication 2314] [LN:Ba arr] [FN:ne- pott] [DE:Ba	PUR OPERON 9685] cillus subtilis] nown] [OR:Bacillus A] ation origin.] SUB0001] gative regulation acillus subtilis
Description sp:[LN:PURR_BACSU] [AC:P37551] [GN:REPRESSOR] [SP:P37551] [DB:swissprote [PN:transcription repressor of puring [DB:pir2] >gp:[GI:d1005824:g467436] subtilis] [SR:Bacillus subtilis (subtilis) [SR:Bacillus subtilis (subtilis) [DE:B. subtilis DE:LE:118041] [RE:118898] [DI:direct] [AC:Z99104:AL009126] [PN:transcripts of the purine operon] [OR:Bacillus subtiles genome (section 1 of 21):	PURR] [O. t] >pir: ne opero: [LN:BAC o_specie NA, 180 >gp:[GI ional resubtilis from 1 to	R:BACII [LN:S60 n purk] 180K] s:Marbo kilobas :e1181: gulato:] [DB:0 0213080	LengthLe [867] [28 LLUS SUBT 5076] [AC [AC:D2618 urg, stra se region 980:g2632 r] [GN:pur genpept-h 0.] [NT:a	FILIS] [DE: C:S66076:D6 of [PN:unk: ain:168) DN: a of replication of [FN:new cott] [DE:Balternate go	PUR OPERON 9685] cillus subtilis] nown] [OR:Bacillus A] ation origin.] SUB0001] gative regulation acillus subtilis
Description sp:[LN:PURR_BACSU] [AC:P37551] [GN:REPRESSOR] [SP:P37551] [DB:swissprote [PN:transcription repressor of puring [DB:pir2] >gp:[GI:d1005824:g467436] subtilis] [SR:Bacillus subtilis (subtilis) [SR:Bacillus subtilis DB:genpept-bct1] [DE:B. subtilis DB [LE:118041] [RE:118898] [DI:direct] [AC:Z99104:AL009126] [PN:transcription of the purine operon] [OR:Bacillus subtilis complete genome (section 1 of 21): [SP:P37551] [LE:54439] [RE:55296] [I	PURR] [O. c.] >pir: ne opero: [LN:BAC o_specie NA, 180 >gp:[GI ional resubtilis from 1 to	R:BACII [LN:S60 n purk] 180K] s:Marbo kilobas :e1181: gulato:] [DB:0 0213080	LengthLe [867 28 LLUS SUBT 6076] [AC [GN:pur [AC:D2618 urg, stra se region 980:g2632 r] [GN:pur genpept-h D.] [NT:a	TILIS] [DE: C:S66076:D6 CR] [OR:Ba S5] [PN:unk ain:168) DN ch of replication of replication of the control of t	PUR OPERON 9685] cillus subtilis] nown] [OR:Bacillus A] ation origin.] SUB0001] gative regulation acillus subtilis ene name: yabI]
Description sp:[LN:PURR_BACSU] [AC:P37551] [GN:IREPRESSOR] [SP:P37551] [DB:swissprote [PN:transcription repressor of puring [DB:pir2] >gp:[GI:d1005824:g467436] subtilis] [SR:Bacillus subtilis (subtilis) [SR:Bacillus subtilis (subtilis) [DE:B. subtilis DR [LE:118041] [RE:118898] [DI:direct] [AC:Z99104:AL009126] [PN:transcripts of the purine operon] [OR:Bacillus subtilis genome (section 1 of 21): [SP:P37551] [LE:54439] [RE:55296] [IN:DRF Name]	PURR] [O]	R:BACII [LN:S66 n purk 180K] s:Marbi kilobas :e11819 gulatos] [DB:9 0213086 t]	LengthLe [867 28 LLUS SUBT 5076] [AC [AC:D2618 1rg, stra se region 980:g2632 r] [GN:pu genpept-h 0.] [NT:a	TILIS] [DE: C:S66076:D6 CR] [OR:Ba B5] [PN:unk ain:168) DN n of replic C:314] [LN:Ba arR] [FN:ne cott] [DE:Ba alternate ge AA angth	PUR OPERON 9685] cillus subtilis] nown] [OR:Bacillus A] ation origin.] SUB0001] gative regulation acillus subtilis ene name: yabI] probability

SPORULATION PROTEIN G] [SP:P28016] [DB:swissprot] >pir:[LN:S18900] [AC:S18900] [PN:spoVG protein] [CL:stage V sporulation protein spoVG] [OR:Bacillus megaterium] [DB:pir2] >gp:[GI:g39656] [LN:BMSPOVG] [AC:X62377] [GN:spoVG] [OR:Bacillus megaterium] [DB:genpept-bct1] [DE:B.megaterium spoVG and tms genes.] [SP:P28016] [LE:31] [RE:321]

[DI:direct]

ORF Name	NTID	AAID		<u>AA</u>	probability	
A17503000980 34001510 c2 218			LengthLer			_
	182	3954	798 26	5 916	6.4e-92	
Description sp:[LN:YABD_BACSU] [AC:P37545] [GN:Y KD PROTEIN IN METS-KSGA INTERGENIC R [AC:S66068:E69738] [PN:conserved hy [CL:hypothetical protein HI0454] [OR >gp:[GI:d1005816:g467428] [LN:BAC180 [SR:Bacillus subtilis (sub_species:M subtilis DNA, 180 kilobase region of [DI:direct] >gp:[GI:e1181972:g263230 [FN:unknown] [OR:Bacillus subtilis] genome (section 1 of 21): from 1 to2 [SP:P37545] [LE:47704] [RE:48471] [D	EGION] potheti :Bacill K] [AC: larburg, replic [6] [LN: [DB:gen 13080.]	(SP:P3 cal pr us sub D26185 strai ation BSUB00 pept-b [NT:s	7545] [DB otein yabl tilis] [Di] [PN:unkr n:168) DNr origin.] 01] [AC:ZS ct1] [DE:H	:swissprot D] [GN:yak B:pir2] nown] [OR: A] [DB:gen [LE:111306 99104:AL00 Bacillus s	Bacillus subtil pept-bctl] [DE: [RE:112073] [P126] [GN:yabD] Subtilis complet	is] B.
ORF Name	NTID	AAID	NT A	AA score	probability	
AI7503000980_34428515_c1_191	183	3955	<u>Bengenber</u> 		6.0e-25	\neg
Description sp:[LN:VEG_BACSU] [AC:P37466] [GN:VE [SP:P37466] [DB:swissprot] >pir:[LN: protein veg] [GN:veg] [OR:Bacillus [LN:BAC180K] [AC:D26185] [PN:unknown subtilis (sub_species:Marburg, strai 180 kilobase region of replication o >gp:[GI:e1181977:g2632311] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-b 1 of 21): from 1 to213080.] [SP:P374	S66073] subtili] [GN:v n:168) rigin.] 001] [A	[AC:Ss] [DBeg] [O DNA] [[LE:1 C:Z991 E:Baci	66073:C697 :pir2] >gr R:Bacillus DB:genpept 16363] [RI 04:AL00912 llus subti	730] [PN: p:[GI:d100 s subtilis c-bct1] [D E:116623] 26] [GN:ve ilis compl	hypothetical [5821:g467433] [] [SR:Bacillus [E:B. subtilis Di [DI:direct] [g] [FN:unknown] [ete genome (sec	
ORF Name	NTID	AAID	NT A	A gth score	probability	
A17503000980_34571011_c3_252	184	3956	1608 539		2.8e-176	7
Description sp:[LN:MFD_BACSU] [AC:P37474] [GN:MFCOUPLING FACTOR (TRCF)] [SP:P37474] [PN:transcription-repair coupling facoupling protein:DEAD/H box helicase psp:[GI:d1005832:g467444] [LN:BAC180 factor] [GN:mfd] [OR:Bacillus subtil strain:168) DNA] [DB:genpept-bct1] [Strain:168) DNA] [DB:genpept-bct1] [Replication origin.] [LE:124030] [RE [LN:BSUB0001] [AC:Z99104:AL009126] [FN:probably involved in homologous [DE:Bacillus subtilis complete genom [LE:60428] [RE:63961] [DI:direct]	[DB:swi ctor mf homolo K] [AC: is] [SR DE:B. s :127563 PN:tran DNA] [O	ssprot d] [GN gy] [O D26185 :Bacil ubtili] [DI: script R:Baci] >pir:[LN:mfd] [CI R:Bacillus] [PN:tran lus subtil s DNA, 180 direct] >g ion-repair llus subti	N:S66085] S:transcri S:subtilis S:cription S:s (sub_s S) kilobase Sp:[GI:e11 S:coupling Llis] [DB:	[AC:S66085:F6969] ption-repair] [DB:pir2] -repair coupling pecies:Marburg, region of 81988:g2632322] factor] [GN:mfogenpept-bct1]	57] g d]
ORF Name	NTID	AAID	NT A	A gth score	probability	
A17503000980 35260887 cl 193	185	13957	1077 146			

ORF Name	NTID	AAID	<u>NT</u> LengthL	AA ength	score	probability
AI7503000980_3553_c3_245	186	3958	972	323	382	2.5e-35
Description	J [·	
sp:[LN:HOLB_BACSU] [AC:P37540] [GN:POLYMERASE III, DELTA' SUBUNIT,] [STEAT [AC:S66061:C69642] [PN:DNA polymerates [GN:holB] [OR:Bacillus subtilis] [IAC:D26185] [PN:similar to B. subtilis [Subtilis (sub_species:Marburg, strates [AS kilobase region of replication of page:[GI:e1181964:g2632298] [LN:BSUB6 (delta' subunit)] [GN:holB] [OR:BacilE:Bacillus subtilis complete genore [NT:alternate gene name: yaaS] [SP:Name [NT:alternate gene name [NT:alternate gene nam	P:P37540 ase III DB:pir2] lis DnaH in:168) origin.] 0001] [A illus su me (sect] [DB: (delta >gp:[] [OR: DNA] [[LE:1 .C:Z991 btilis	swisspro GI:d1005 Bacillus DB:genpe .04265] [.04:AL009 c] [DB:geof 21):	ot] >psit) holdstand holds	ir:[LN lB:dna 467421 ilis] [1] [[5254] [PN:DN -bct1] L to21	N:S66061] AH homolog holB] L] [LN:BAC180K] [SR:Bacillus DE:B. subtilis DNA, [DI:direct] NA polymerase III [EC:2.7.7.7]
ORF Name	NTID	AAID	<u>NT</u> LengthL	AA ength	score	probability
AI7503000980_3916087_c2_216	187	3959	261	36	243	1.3e-20
Description		·				
pir:[LN:A69742] [AC:A69742] [PN:con [CL:hypothetical protein 312] [OR:Ba pgp:[GI:e1181968:g2632302] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-lof 21): from 1 to213080.] [NT:sime [DI:direct]	acillus 0001] [A bctl] [D	subtil C:Z991 E:Baci	is] [DB: 04:AL009 llus sub	pir2] 9126] otilis	[GN:ya	zA] [FN:unknown] ete genome (section
ORF Name	NTID	AAID	<u>NT</u> Length <u>L</u>	<u>AA</u> ength	score	probability
A17503000980_3937950_£2_71	188	3960		51		
Description						
TIH-OV						
ORF Name	NTID	AAID	NT LengthL	AA ength	score	probability
A17503000980_3938838_c1_210	189	3961	846 2	281	1087	4.8e-110
Description						
sp:[LN:DHPS_STAHA] [AC:Q59919] [GN:FIDE:PYROPHOSPHORYLASE) (DHPS)] [SP:QILN:SHU40768] [AC:U40768] [PN:dihydraemolyticus] [DB:genpept-bct2] [EC:synthase A (cysK) anddihydroneopterianddihydropteroate synthase (folp) [EC:Polymore anddihydropteroate synthase (folp) [EN:Polymore and [EN:Polymore	Q59919] ropteroa :2.5.1.1 in aldol	[DB:sw te syn 5] [DE ase (f	issprot] thase] [:Staphyl olQ) gen	>gp: GN:fol ococcu es, pa	[GI:gl .P] [O is hae irtial	118002] R:Staphylococcus molyticus cysteine
[DI:direct]	gene, co	mplete	cds.] [NT:DHE	s] [L	E:692] [RE:1495]
	NTID	Mplete AAID	NT LengthL	<u>AA</u>	core	E:692] [RE:1495] probability

ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000980 4723192_c1 196	191	3963	
Description			
sp:[LN:SP5C_BACSU] [AC:P37470] [GN:S [DE:SPORULATION PROTEIN C)] [SP:P374 [AC:C69715:S66083] [PN:stage V spor spoVC] [GN:spoVC] [CL:peptidyl-tRNA	70] [DB ulation hydrology [AC: subtilication	:swissp protei ase] [O D26185] s (sub_ kilobas :e11819 s subti from 1	orot] >pir:[LN:C69715] In spoVC:spore coat formation protein OR:Bacillus subtilis] [DB:pir2] [PN:stage V sporulation] [GN:spoVC] species:Marburg, strain:168) DNA] se region of replication origin.] O86:g2632320] [LN:BSUB0001] Llis] [DB:genpept-bct1] [DE:Bacillus Lto213080.] [NT:thermosensitive
ORF Name	NTID	<u>AAID</u>	NT AA score probability
A17503000980_4775312_c3_257	192	3964	405 134 436 4.7e-41
Description		<u> </u>	
subtilis] [SR:Bacillus subtilis (sub [DB:genpept-bct1] [DE:B. subtilis DN [LE:133226] [RE:133612] [DI:direct] [PN:unknown] [OR:Bacillus subtilis] division protein (divIC) gene, compl	C REGION 69740] divIC 3 ansfera [LN:BAC species A, 180] >gp:[GI [DB:gen] ete cds end of] [DI:d: GN:yabR lis com	N] [SP: [PN:po '-regio se homo 180K] [s:Marbu kilobas :g38517 pept-bc ,transf cds.] irect]] [FN:u plete g	P37560] [DB:swissprot] plyribonucleotide on hypothetical protein] [GN:yabR] plog yabR] [OR:Bacillus subtilis] [AC:D26185] [PN:unknown] [OR:Bacillus arg, strain:168) DNA] se region of replication origin.] [AB] [LN:BACDIVIC] [AC:L23497] [AT] [DE:Bacillus subtilis cell are RNA genes, complete sequence and [NT:orf128; homologous to RNA binding origins or specific section of a subtilis are specific section or subtilis or subtilis or specific section or subtilis or subtilis or specific section or subtilis or
ORF Name	NTID	AAID I	NT AA LengthLength score probability
A17503000980_4884625_c1_189	193	3965	549 182 445 5.2e-42
Description sp:[LN:YABF_BACSU] [AC:P37547] [GN:Y. KD PROTEIN IN METS-KSGA INTERGENIC R [AC:S66070:G69738] [PN:conserved hysubtilis] [DB:pir2] >gp:[GI:d1005818 [OR:Bacillus subtilis] [SR:Bacillus [DB:genpept-bct1] [DE:B. subtilis DN. [LE:113687] [RE:114247] [DI:direct] [AC:Z99104:AL009126] [GN:yabF] [FN:um [DE:Bacillus subtilis complete genome to hypothetical proteins] [SP:P37547]	EGION] pothetic :g467430 subtilis A, 180 } >gp:[GI: nknown] e (sect	[SP:P37 cal pro 0] [LN: s (sub_ kilobas: :ell819 [OR:Ba ion 1 o	547] [DB:swissprot] >pir:[LN:S66070] tein yabF] [GN:yabF] [OR:Bacillus BAC180K] [AC:D26185] [PN:unknown] species:Marburg, strain:168) DNA] e region of replication origin.] 74:g2632308] [LN:BSUB0001] cillus subtilis] [DB:genpept-bct1] f 21): from 1 to213080.] [NT:similar
ORF Name	NTID	AAID I	NT AA LengthLength
AT7503000980 5317151 c3 264	17 94	_	177 EQ

ORF Name

NTID AAID I PORT NEORT NEO

ID AAID LengthLength score probability

8.0e-85

Description

sp:[LN:YABN_BACSU] [AC:P37556] [GN:YABN] [OR:BACILLUS SUBTILIS] [DE:HYPOTHETICAL 56.1 KD PROTEIN IN MFD-DIVIC INTERGENIC REGION] [SP:P37556] [DB:swissprot] >pir:[LN:S66088] [AC:S66088:E69739] [PN:conserved hypothetical protein yabN:beta-lactamase regulatory protein homolog yabN] [GN:yabN] [CL:beta-lactamase regulatory protein homolog: beta-lactamase regulatory protein homology] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:d1005835:g467447] [LN:BAC180K] [AC:D26185] [PN:unknown] [OR:Bacillus subtilis] [SR:Bacillus subtilis (sub_species:Marburg, strain:168) DNA] [DB:genpept-bct1] [DE:B. subtilis DNA, 180 kilobase region of replication origin.] [LE:130005] [RE:131474] [DI:direct] >gp:[GI:e1181991:g2632325] [LN:BSUB0001] [AC:Z99104:AL009126] [GN:yabN] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 1 of 21): from 1 to213080.] [NT:similar to hypothetical proteins] [SP:P37556] [LE:66403] [RE:67872] [DI:direct]

 ORF Name
 NTID
 AAID
 NT AA Length Length
 score
 probability

 A17503000980_5367813_c3_244
 196
 3968
 543
 180
 432
 1.2e-40

Description

sp:[LN:ATDA ECOLI] [AC:P37354] [GN:SPEG] [OR:ESCHERICHIA COLI] [EC:2.3.1.57] [DE:ACETYLTRANSFERASE) (SAT)] [SP:P37354] [DB:swissprot] >gp:[GI:d1016007:g1742583] [LN:D90799] [AC:D90799:AB001340] [PN:Spermidine N1-acetyltransferase (EC 2.3.1.57)] [GN:speG] [OR:Escherichia coli] [SR:Escherichia coli (strain:K12) DNA, clone lib:Kohara lambda minise] [DB:genpept-bct1] [DE:E.coli genomic DNA, Kohara clone #308(35.3-35.7 min.).] [NT:ORF_ID:o309#16; similar to [SwissProt Accession] [LE:<18154] [RE:18711] [DI:direct] >gp:[GI:d1016029:g1742606] [LN:D90800] [AC:D90800:AB001340] [PN:Spermidine N1-acetyltransferase (EC 2.3.1.57)] [GN:speG] [OR:Escherichia coli] [SR:Escherichia coli (strain:K12) DNA, clone_lib:Kohara lambda minise] [DB:genpept-bct1] [DE:E.coli genomic DNA, Kohara clone #309(35.4-35.7 min.).] [NT:ORF ID:o309#16; similar to [SwissProt Accession] [LE:<14930] [RE:15487] [DI:direct] >gp:[GI:d1016031:g1742609] [LN:D90801] [AC:D90801:AB001340] [PN:Spermidine N1-acetyltransferase (EC 2.3.1.57)] [GN:speG] [OR:Escherichia coli] [SR:Escherichia coli (strain:K12) DNA, clone lib:Kohara lambda minise] [DB:genpept-bct1] [DE:E.coli genomic DNA, Kohara clone #310(35.7-36.0 min.).] [NT:ORF_ID:o309#16; similar to [SwissProt Accession] [LE:<814] [RE:1371] [DI:direct]

 ORF Name
 NTID
 AAID
 NTID
 AAID
 NTID
 LengthLength
 score
 probability

 A17503000980_581260_f3_137
 197
 3969
 183
 60
 183
 60
 NO-HIT
 NO-HIT</t

ORF Name NTID AAID Length score probability

Description

sp:[LN:YABH_BACSU] [AC:P37550] [GN:YABH] [OR:BACILLUS SUBTILIS] [DE:HYPOTHETICAL 31.7
KD PROTEIN IN SSPF-PURR INTERGENIC REGION (ORF1)] [SP:P37550] [DB:swissprot]
>pir:[LN:S66075] [AC:S66075:A69739] [PN:conserved hypothetical protein yabH] [GN:yabH]
] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:d1005823:g467435] [LN:BAC180K] [AC:D26185]
[PN:unknown] [OR:Bacillus subtilis] [SR:Bacillus subtilis (sub_species:Marburg,
strain:168) DNA] [DB:genpept-bct1] [DE:B. subtilis DNA, 180 kilobase region of
replication origin.] [LE:117116] [RE:117985] [DI:direct] >gp:[GI:e1181979:g2632313]
[LN:BSUB0001] [AC:Z99104:AL009126] [GN:yabH] [FN:unknown] [OR:Bacillus subtilis]
[DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 1 of 21): from 1
to213080.] [NT:similar to hypothetical proteins] [SP:P37550] [LE:53514] [RE:54383]
[DI:direct]

 ORF Name
 NTID
 AAID
 NT AAID
 AAID
 score
 probability

 A17503000980_6136562_c2_223
 199
 3971
 402
 133
 361
 4.1e-33

Description

sp:[LN:YABJ_BACSU] [AC:P37552] [GN:YABJ] [OR:BACILLUS SUBTILIS] [DE:HYPOTHETICAL 13.7
KD PROTEIN IN PURR-SPOVG INTERGENIC REGION (ORF2)] [SP:P37552] [DB:swissprot]
>pir:[LN:S66077] [AC:S66077:B69739] [PN:conserved hypothetical protein yabJ] [GN:yabJ]
[CL:hypothetical protein HI0719] [OR:Bacillus subtilis] [DB:pir2]
>gp:[GI:d1005825:g467437] [LN:BAC180K] [AC:D26185] [PN:unknown] [OR:Bacillus subtilis]
[SR:Bacillus subtilis (sub_species:Marburg, strain:168) DNA] [DB:genpept-bct1] [DE:B.
subtilis DNA, 180 kilobase region of replication origin.] [LE:118895] [RE:119272]
[DI:direct] >gp:[GI:e1181981:g2632315] [LN:BSUB0001] [AC:Z99104:AL009126] [GN:yabJ]
[FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete
genome (section 1 of 21): from 1 to213080.] [NT:similar to hypothetical proteins]
[SP:P37552] [LE:55293] [RE:55670] [DI:direct]

 ORF Name
 NTID
 AAID
 NT Length Length
 Score
 probability

 A17503000980_6742943_c2_217
 200
 3972
 1983
 660
 2292
 9.8e-238

<u>Description</u>

sp:[LN:SYM_BACSU] [AC:P37465] [GN:METS] [OR:BACILLUS SUBTILIS] [EC:6.1.1.10]
[DE:(METRS)] [SP:P37465] [DB:swissprot] >pir:[LN:S66067] [AC:S66067:E69657]
[PN:methionine--tRNA ligase, metS:methionyl-tRNA synthetase metS] [GN:metS]
[CL:methionine--tRNA ligase] [OR:Bacillus subtilis] [EC:6.1.1.10] [DB:pir2]
>gp:[GI:d1005815:g467427] [LN:BAC180K] [AC:D26185] [PN:methionyl-tRNA synthetase]
[GN:metS] [OR:Bacillus subtilis] [SR:Bacillus subtilis (sub_species:Marburg,
strain:168) DNA] [DB:genpept-bct1] [DE:B. subtilis DNA, 180 kilobase region of
replication origin.] [LE:109233] [RE:111227] [DI:direct] >gp:[GI:e1181971:g2632305]
[LN:BSUB0001] [AC:Z99104:AL009126] [PN:methionyl-tRNA synthetase] [GN:metS]
[OR:Bacillus subtilis] [DB:genpept-bct1] [EC:6.1.1.10] [DE:Bacillus subtilis complete
genome (section 1 of 21): from 1 to213080.] [SP:P37465] [LE:45631] [RE:47625]
[DI:direct]

ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	score	probability
AI7503000980_6834427_c2_227	201	3973		49	93	0.00010
Description	L		الـــــــا لــ		<u> </u>	
sp:[LN:YABO_BACSU] [AC:P37557] [GN:YPROTEIN IN MFD-DIVIC INTERGENIC REGION [AC:S66089:F69739] [PN:conserved hysubtilis] [DB:pir2] >gp:[GI:d1005836] [OR:Bacillus subtilis] [SR:Bacillus [DB:genpept-bct1] [DE:B. subtilis DN:[LE:131477] [RE:131737] [DI:direct] [AC:Z99104:AL009126] [GN:yabO] [FN:upper [DE:Bacillus subtilis complete genome to hypothetical proteins] [SP:P37557]	ON] [SP: pothetic :g467448 subtilis A, 180 k >gp:[GI: nknown] e (secti	P3755 cal pro B] [LN s (sub cilobas e1181; [OR:Ba	7] [DB: otein y :BAC180 _specie se regi 992:g26 acillus of 21):	wissp vabO] (OK] [AC es:Mark on of [32326] s subti	orot] : [GN:yal] C:D2618 ourg, : replic [LN:I .lis]	ppir:[LN:S66089] OO] [OR:Bacillus BS] [PN:unknown] Strain:168) DNA] Cation origin.] BSUB0001] [DB:genpept-bct1] L3080.] [NT:similar
ORF Name	NTID	AAID	NT Length			probability
A17503000980_7036526_c1_212	202	3974	159	52	86	0.00057
Description pir:[LN:C64571] [AC:C64571] [PN:hype [DB:pir2] >gp:[GI:g2313526] [LN:AE00] predicted coding region HP0411] [GN:] [DB:genpept-bct2] [DE:Helicobacter progenome.] [NT:hypothetical protein; in [DI:direct]	0557] [A HP0411] ylori 26	C:AE00 OR:H6 695 se	00557:A elicoba ection	E00051 cter p 35 of	.1] [PN ylori 134 of	N:H. pylori 26695] T the complete
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
AI7503000980_761_c1_187	203	3975		249		3.1e-92
Description		L			l L	
gp:[GI:e1386911:g4454321] [LN:SAU1328 [OR:Staphylococcus aureus] [DB:genpe (partial).] [NT:ORF1] [LE:434] [RE:1	pt-bct1]	[DE:	Staphyl			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
AI7503000980_829800_c1_201	204	3976	1575	524	700	4.9e-69
Description gp:[GI:g4090864] [LN:AF023181] [AC:Al [GN:ltrB] [OR:Listeria monocytogenes] transcription-repair coupling factor (ltrB), and DivIC homolog(divL) genes	DB:ge (mfdL),	npept-	-bct2] emperat	[DE:Li ure re	steria quirem	monocytogenes ment B protein
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000980_976638_c1_183	205	3977	1371	456	556	9.0e-54
Description						
sp:[LN:YAAO_BACSU] [AC:P37536] [GN:YAKD PROTEIN IN XPAC-ABRB INTERGENIC RI [AC:S66057:F69737] [PN:lysine decarboutilis] [DB:pir2] >gp:[GI:d1005805] lysine decarboxylase] [OR:Bacillus su (sub_species:Marburg, strain:168) DNA kilobase region of replication origin >gp:[GI:e1181960:g2632294] [LN:BSUB00 [OR:Bacillus subtilis] [DB:genpept-bot of 21): from 1 to213080.] [NT:simin [RE:39160] [DI:direct]	EGION] [boxylase :g467417 ubtilis] A] [DB:g n.] [LE: 001] [AC ct1] [DE	SP:P37 homo] [LN: [SR:Fenpept 101320 :Z9910	7536] [: Log yaa: :BAC180. Bacillu: :-bct1] D] [RE: D4:AL00.	DB:swi O] [GN K] [AC s subt [DE:B 102762 9126] btilis	ssprot :yaaO :D2618 ilis . subt] [DI: [GN:ya compl] >pir:[LN:S66057]] [OR:Bacillus 5] [PN:similar to ilis DNA, 180 direct] a0] [FN:unknown] ete genome (section

ORF Name	NTID	AAID LengthLength score probability
A17503000980_9876005_£3_159	206	3978 126 41
Description		
NO-HIT		
ORF Name	NTID	AAID <u>NT AA</u> LengthLength score probability
AI7503000981_10392555_f3_17	207	3979 180 59
Description		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000981_134392_t3_23	208	3980 153 50
Description		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
AI7503000981_1385927_f3_22	209	3981 369 122 85 0.039
Description		
	falcipa:	
ORF Name	NTID	AAID NT AA score probability
AI7503000981_14882135_f2_11	210	3982 2112 703 167 1.6e-09
REGULATORY PROTEIN] [SP:P10411] [DB: [AC:A29625:S56347:E65221] [PN:melib [CL:arabinose operon regulatory prot >gp:[GI:g536963] [LN:ECOUW93] [AC:U1 [OR:Escherichia coli] [DB:genpept-bc from 92.8 to 00.1 minutes.] [NT:CG S [DI:complement] >gp:[GI:g1790559] [L melibiose operon] [GN:melR] [FN:regulation of the second content of the	swissproiose operation of the No. N:AE0004 lator; I	eron regulatory protein] [GN:melR] R:Escherichia coli] [DB:pir1] [MP:93 min] GN:melR] [FN:regulatory gene] :Escherichia coli K-12 chromosomal region 18166] [LE:31548] [RE:32456] 484] [AC:AE000484:U00096] [PN:regulator of Degradation of small molecules:] :Escherichia coli K-12 MG1655 section 374 of
ORF Name AI7503000981_23634578_f3_18 Description	<u>NTID</u>	AAID NT AA Score probability 3983 150 49
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000981_23860887_f2_13	212	3984 141 46
Description NO-HIT		

ORF Name	NTID	AAID	NT AA score	e probability
AI7503000981_25634627_£2_12	213	3985		1.4e-25
Description	لــــا ك	<u> </u>	J	
sp:[LN:GS26_BACSU] [AC:P80238:P9658 STRESS PROTEIN 26 (GSP26)] [SP:P802 [AC:G69768] [PN:general stress pro [DB:pir2] >gp:[GI:d1020013:g1881233 subtilis] [SR:Bacillus subtilis (st subtilis genome sequence, 148 kb se [NT:FUNCTION UNKNOWN.] [LE:6862] [R [LN:BSUB0003] [AC:Z99106:AL009126] [DB:genpept-bct1] [DE:Bacillus subt to611850.] [NT:alternate gene name: [RE:71028] [DI:direct]	38:P9658 tein hom] [LN:AB rain:168 quence o E:7284] [GN:ydaG ilis com	o) [DB olog y 001488) DNA] f the [DI:di	:swissprot] >pir: daG] [GN:ydaG] [G] [AC:AB001488] [G [DB:genpept-bct1] regionbetween 35 a rect] >gp:[GI:e118 unknown] [OR:Baci] genome (section 3	[LN:G69768] DR:Bacillus subtilis] GN:ydaG] [OR:Bacillus] [DE:Bacillus] and 47 degree.] B2388:g2632722] Ilus subtilis] of 21): from 402751
ORF Name	NTID	AAID	NT AA LengthLength	probability
A17503000981 2584538 c1 29	214	3986	7 132 43	
Description	J LJ	L	J	
NO-HIT				
ORF Name	NTID	AAID	NT AA score	e probability
A17503000981_29352342_c2_36	215	3987	1296 431 1218	6.3e-124
Description	ــــــا ا	L	J	
sp:[LN:GLTT_BACST] [AC:P24943] [GN:GSP:P24943] [DB:swissprot] >pir:[LN transport protein:sodium/proton-glu stearothermophilus] [DB:pir2] >gp:[Gutamate symport protein] [GN:gltP stearothermophilus (library: ATCC 7 stearothermophilus proton glutamate [RE:1375] [DI:direct]	:S26247] tamate s GI:g1430] [OR:Ba 954) DNA	[AC:S ymport 00] [L cillus] [DB:	26247] [PN:glutam protein] [GN:gltF N:BACGLTPA] [AC:M8 stearothermophilu genpept-bct1] [DE:	nate/aspartate P] [OR:Bacillus B6508] [PN:proton as] [SR:Bacillus Bacillus
ORF Name	NTID	AAID	NT AA Length Length	probability
A17503000981_30081465_c2_37	216	3988	147 48	
Description NO-HIT	J []			
ORF Name	NTID	AAID	NT AA LengthLength	probability
A17503000981_32242200_f1_2	217	3989	378 125	
Description	J []		J	
NO-HIT				
ORF Name	NTID	AAID	NT AA LengthLength	probability
AI7503000981_3396042_c1_34 Description	218	3990	315 104	

ORF Name AI7503000981_34179828_c2_35 Description NO-HIT	<u>NTID</u> 219	<u>AAID</u> 3991	NT AA LengthLength score	probability
ORF Name AT7503000981_4100336_f2_10 Description NO-HIT	<u>NTID</u> 220	<u>AAID</u> 3992	NT AA LengthLength score	probability
ORF Name A17503000981_6511652_c3_51 Description NO-HIT	NTID 221	<u>AAID</u> 3993	NT AA LengthLength score	probability
ORF Name A17503000981_822786_c1_33 Description NO-HIT	NTID	<u>AAID</u> 3994	NT AA score LengthLength 132 43	probability
ORF Name AI7503000982_10031712_c1_1057 Description sp:[LN:YQJA_BACSU] [AC:P54538] [GN:Y KD PROTEIN IN BMRU-ANSR INTERGENIC R [AC:A69963] [PN:hypothetical protei >gp:[GI:d1013287:g1303952] [LN:BACJH subtilis] [SR:Bacillus subtilis (str [DE:Bacillus subtilis DNA, 283 Kb re [RE:230115] [DI:direct] >gp:[GI:e118 [GN:yqjA] [FN:unknown] [OR:Bacillus complete genome (section 13 of 21): [RE:94472] [DI:complement]	EGION] n yqjA] 642] [A ain:JH6 gion co 5663:g2 subtili	[SP:P5 [GN:y C:D844 42(trp ntaini 634829 s] [DB	LLUS SUBTILIS] [DE:164538] [DB:swissprot] [QjA] [OR:Bacillus 1632:D82370] [PN:YqjA] [C2 PheA1)) DNA] [DB [Dg skin element.] [16] [LN:BSUB0013] [AC [S:genpept-bct1] [DE:16]	<pre> ->pir:[LN:A69963] subtilis] [DB:pir2] [OR:Bacillus :genpept-bct1] LE:229147] :Z99116:AL009126] Bacillus subtilis</pre>

NT AΑ ORF Name NTID AAID score probability LengthLength 3996 AI7503000982 10312561 c3 1455 1236 411 224 996 2.1e-100 Description sp:[LN:RS1H BACSU] [AC:P38494] [GN:YPFD:JOFD] [OR:BACILLUS SUBTILIS] [DE:30S RIBOSOMAL PROTEIN S1 HOMOLOG] [SP:P38494] [DB:swissprot] >pir:[LN:B69935] [AC:B69935] [PN:ribosomal protein S1 homolog homolog ypfD] [GN:ypfD] [CL:Synechocystis ribosomal protein S1] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:g533106] [LN:BSU11687] [AC:U11687] [PN:unknown] [GN:jofD] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis 168 jofA, jofB, MssA homolog (jofC) and ribosomalprotein S1 homolog (jofD) genes, complete cds, and joeB gene, partial cds.] [NT: similar to the Escherichia coli S1 ribosomal] [LE:2651] [RE:3799] [DI:direct] >qp:[GI:e1183733:q2634706] [LN:BSUB0012] [AC:Z99115:AL009126] [GN:ypfD] [FN:unknown] [OR:Bacillus subtilis] [DB:qenpept-bct1] [DE:Bacillus subtilis complete genome (section 12 of 21): from 2195541to 2409220.] [NT:alternate gene name: jofD; similar to ribosomal] [SP:P38494] [LE:198361] [RE:199509] [DI:complement] >gp:[GI:g1146215] [LN:BACSERA] [AC:L47648] [GN:ypfD] [OR:Bacillus subtilis] [DB:qenpept-bct2] [DE:Bacillus subtilis phosphoglycerate dehydrogenase (serA), ypaA, ferredoxin (fer), ypbB, recS, ypbD, ypbE, ypbF, ypbG, ypbH, glutamate dehydrogenase (ypcA), ypdA, ypdB, ypdC, spore cortexlytic enzyme (sleB), ypeB, ypfA, ypfB, cytidine monophosphatekinase (cmk), ypfD, ypgA, yphA, yphB, yphC, NAD+ dependentglycerol-3-phosphate dehydrogenase (glyc), yphE and yphF genes,complete cds.] [NT:similar to Escherichia coli S1 ribosomal protein;] [LE:16888] [RE:18036] [DI:direct] NT AΑ Length Length score ORF Name NTID AAID probability AI7503000982 10334752 f1 178 225 3997 192 63 Description NO-HIT NT AA ORF Name NTID score AAID probability LengthLength AI7503000982 10588877 £3 947 226 3998 162 Description NO-HIT NT AΑ score ORF Name NTID AAID probability

227

NTID

228

3999

AAID

4000

AI7503000982 10635918 c2 1290

A17503000982 10642180 c3 1513

Description NO-HIT

ORF Name

Description NO-HIT

LengthLength

LengthLength

61

40

AΑ

<u>score</u>

probability

123

NT

186

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000982_10666068_c2_1228	229	4001	330	109	196	1.3e-15
Description sp:[LN:CMG3_BACSU] [AC:P25955] [GN:CGPROTEIN 3 PRECURSOR] [SP:P25955] [DB [AC:D30338:A35133:D69603] [PN:exoger 3] [GN:comGC] [OR:Bacillus subtilisted [AC:M29691:M22854] [GN:comG3] [OR:Bacillus subtilisted [Clone: pED4) DNA] [DB:genpept-bct1] comG-(1,2,3,4,5,6,and 7) proteins incomG-(1,2,3,4,5,6,and 7) pr	:swissp nous DN] [DB:p cillus [DE:Ba comG op 9] [LN: subtili lis DNA >gp:[GI exogeno llus su	rot] > A-bind ir2] > subtil cillus eron, BACJH6 s (str., 283 : e1185 us DNA btilis	pir:[LN ing pro gp:[GI: is] [SF subtil complet 42] [AC ain:JH6 Kb regi 739:g26 -bindin comple	N:D3033 Dtein of E:Bacillis (cl.) Lis (cl.) Lis (cl.) Ee cds C:D8443 542(trp.) Lion con 534905] Ing (concete gen	comGC: comGC: llus s lone p lone p lone p comBallian lus s lus s	comG operon protein N:BACCOMGA] ubtilis (strain 168) ED4) :3405] [RE:3701] 370] [PN:ComGC] eA1)) DNA] ng skin element.] BSUB0013] ce)] [OR:Bacillus section 13 of 21):
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
AY7503000982_1070437_c3_1482	230	4002		297		6.8e-72
Description sp:[LN:YPCP_BACSU] [AC:P54161] [GN:YI [DE:POTENTIAL 5'-3' EXONUCLEASE,] [SI [AC:H69933] [PN:5'-3' exonuclease ha [DB:pir2] >gp:[GI:g1256623] [LN:BACYI [GN:ypcP] [OR:Bacillus subtilis] [DB clone) DNA region between the serA and si 5' end of the] [LE:9565] [RE:10455] [LN:BSUB0012] [AC:Z99115:AL009126] [OB:genpept-bct1] [DE:Bacillus subtilis] 2195541to 2409220.] [NT:similar to 5 [RE:115587] [DI:complement]	P:P5416 omolog ACA] [A :genpep ndkdg l [DI:dir GN:ypcP lis com	1] [DB ypcP] C:L772 t-bct1 oci.] ect] >] [FN: plete	:swissp [GN:ypc 46] [PN] [DE:E [NT:36. gp:[GI: unknown genome	orot] : CP] [C N:exode Bacillu 1% ide :e11836 n] [OR:	pir:[] OR:Bac eoxyril is sub entity 547:g20 :Bacil ion 12	LN:H69933] illus subtilis] bonuclease] tilis (YAC10-9 with 219 aa at the 634620] lus subtilis] of 21): from
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000982 10938927 f2 492	231	4003		43	: 1	
Description			لــــاد		J	
NO-HIT						
ORF Name A17503000982_10969050_c2_1193	NTID	<u>AAID</u> 4004	<u>NT</u> Length			probability 3.5e-187
Description sp:[LN:LEPA_BACSU] [AC:P37949] [GN:LI LEPA] [SP:P37949] [DB:swissprot] >pin lepA] [GN:lepA] [CL:GTP-binding memble homology] [OR:Bacillus subtilis] [DB [AC:D84432:D82370] [PN:YqeQ] [OR:Bacillus subtilis] [DB [ac:D84432:D82370] [PN:YqeQ] [OR:Bacillus subtilis] [DB [strain:JH642(trpC2 PheA1)) DNA] [DB region containing skin element.] [LE >gp:[GI:e200671:g1122398] [LN:BSLEPON [DB:genpept-bct1] [DE:B.subtilis lepx [DI:direct] >gp:[GI:e1183781:g2634997] [PN:GTP-binding protein] [GN:lepA] [GDE:Bacillus subtilis complete genome [NT:alternate gene name: yqxB, yqeQ]	r:[LN:G brane p :pir2] illus s :genpep :86866] RF] [AC A and h 7] [LN: DR:Baci e (sect	69649] rotein >gp:[G ubtili t-bct1 [RE:8 :X9165 emN ge: BSUB00 llus si	[AC:G6 lepA:t I:d1013 s] [SR:] [DE:B 8704] [5] [GN: nes.] [14] [AC ubtilis of 21)	ransla :ransla :139:gl Bacillu :Bacillu :DI:dir :lepA] :SP:P37 ::Z9911 :: from	[PN:0 ation 6 .303804 .us subtect] [OR:Ba [949] .7:AL00 genper	GTP-binding protein elongation factor Tu 4] [LN:BACJH642] otilis cilis DNA, 283 Kb acillus subtilis] [LE:128] [RE:1966] 09126] ot-bct1]

ORF Name	NTID	AAID NT AA score probability
A17503000982_11194067_c3_1378	233	4005 747 248 431 1.6e-40
Description sp:[LN:YQEM_BACSU] [AC:P54458] [GN:Y KD PROTEIN IN AROD-COMER INTERGENIC [AC:A69952] [PN:conserved hypotheti [OR:Bacillus subtilis] [DB:pir2] >gp [AC:D84432:D82370] [PN:YqeM] [OR:Bacc (strain:JH642(trpC2 PheA1)) DNA] [DB region containing skin element.] [LE >gp:[GI:el183791:g2635007] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-b 14 of 21): from 2599451to 2812870.] [RE:42660] [DI:complement] ORF Name	QEM] [O] REGION] cal proficillus standard standa	R:BACILLUS SUBTILIS] [DE:HYPOTHETICAL 28.3 [SP:P54458] [DB:swissprot] >pir:[LN:A69952] tein yqeM] [GN:yqeM] [CL:bioC homology] 013129:g1303794] [LN:BACJH642] ubtilis] [SR:Bacillus subtilis t-bct1] [DE:Bacillus subtilis DNA, 283 Kb [RE:77513] [DI:direct] C:Z99117:AL009126] [GN:yqeM] [FN:unknown] E:Bacillus subtilis complete genome (section ilar to hypothetical proteins] [LE:41917] AAID NT AA LengthLength score probability
A17503000982_11203763_c1_985	234	4006 132 43
Description NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000982_11881313_f1_285	235	4007 174 57
Description		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000982_12116562_f2_354	236	4008 123 40
Description NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000982_1218750_c1_996	237	4009 681 226 336 1.8e-30
	npept-bo] [PN:competence protein] [GN:celA] ct2] [DE:Streptococcus pneumoniae competence genes, complete cds; and unknown gene.]
ORF Name	NTID	AAID NT AA score probability LengthLength
A17503000982_1226553_c2_1270	238	4010 141 46
Description NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000982_1229750_c3_1419	239	4011 468 155 505 2.3e-48
Description		
	OR:Bacil	C:AJ010954] [PN:arginine repressor] llus stearothermophilus] [DB:genpept-bct1] nd partial recN gene.] [LE:196] [RE:645]

ORF Name NTID AAID NT AA score probability
AITD AAID LengthLength Score probability AI7503000982_1345752_f3_780 Description Sp: [LN:YQKF_BACSU] [AC:P54569] [GN:YQKF] [OR:BACILLUS SUBTILIS] [DE:HYPOTHETICAL DXIDOREDUCTASE IN ANSR-BMRU INTERGENIC REGION] [SP:P54569] [DB:swissprot] Spir: [LN:H69966] [AC:H69966] [PN:conserved hypothetical protein yqkF] [GN:yqkF] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:d1013318:g1303983] [LN:BACJH642] [AC:D84432:D82370] [PN:YqkF] [OR:Bacillus subtilis] [SR:Bacillus subtilis (strain:JH642(trpC2 PheA1)) DNA] [DB:genpept-bct1] [DE:Bacillus subtilis DNA, 283 Kb region containing skin element.] [LE:259404] [RE:260324] [DI:complement] Spp:[GI:e1185631:g2634797] [LN:BSUB0013] [AC:Z99116:AL009126] [GN:yqkF] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section
13 of 21): from 2395261to 2613730.] [NT:similar to hypothetical proteins] [SP:P54569] [LE:63295] [RE:64215] [DI:direct]
NTID AAID NTID AAID LengthLength score probability A17503000982_1359450_f2_544 Description NO-HIT
NTID AAID NTID AAID LengthLength Score probability A17503000982_136068_f1_227 Description Oir: [LN:D70585] [AC:D70585] [PN:probable glyS protein] [GN:glyS] [CL:Mycoplasma genitalium glycinetRNA ligase] [OR:Mycobacterium tuberculosis] [DB:pir2] Ogg: [GI:e315164:g2078043] [LN:MTCY27] [AC:Z95208:AL123456] [PN:glyS] [GN:glyS] [OR:Mycobacterium tuberculosis] [DB:genpept-bct1] [DE:Mycobacterium tuberculosis H37Rv complete genome; segment 104/162.] [NT:Rv2357c, (MTCY27.23-MTCY98.26), len: 463 aa. glyS,] [LE:19] [RE:1410] [DI:complement]
NTID AAID <u>NT AA</u> score probability A17503000982_13674130_f3_703 243
DRF Name NTID AAID NT AA Score probability A17503000982 13853500 f2 374 [244 [4016 [246]81]

ORF Name AAID

probability

AI7503000982 13876943 cl 1122

245

4017 1488

2.6e-88

Description

pir:[LN:B69610] [AC:B69610:JC5744] [PN:carboxy-terminal processing proteinase ctpA,:tail-specific endopeptidase Prc] [GN:ctpA] [CL:carboxyl-terminal processing proteinase] [OR:Bacillus subtilis] [EC:3.4.99.-] [DB:pir2] >gp:[GI:g2529476] [LN:AF006665] [AC:AF006665] [PN:OrfRM1] [GN:orfRM1] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis 168 region at 182 min containing the cge genecluster.] [NT:similar to the E. coli Prc and carboxyl-terminal] [LE:22886] [RE:24286] [DI:direct] >gp:[GI:g2415395] [LN:AF015775] [AC:AF015775] [PN:proteinase] [GN:ctpA] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis YodA (yodA), YodB (yodB), YodC (yodC), YodD(yodD), ABC-transporter (yodE), permease (yodF), proteinase (ctpA), YodH (yodH), YodI (yodI), carboxypeptidase (yodJ), purinenucleoside phosphorylase (deoD), YodL (yodL), YodM (yodM), YodN(yodN), YodO (yodO), YodP (yodP), acetylornitine deacetylase(argE), butirate-acetoacetate CoA transferase (yodR), butyrateacetoacetate-CoA transferase (yodS), YodT (yodT), CgeE (cgeE), CgeD(cgeD), CgeC (cgeC), CgeA (cgeA), CgeB (cgeB), YzxA (yzxA), UDP-glucose epimerase (yodU), YodV (yodV), and YodW (yodW) genes, complete cds; and YodZ (yodZ) gene, partial cds.] [NT:similar to Synechocystis sp. PCC6803] [LE:5846] [RE:7246] [DI:complement] >gp:[GI:e1185430:g2634351] [LN:BSUB0011] [AC:Z99114:AL009126] [PN:carboxy-terminal processing protease] [GN:ctpA] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 11 of 21): from 2000171to 2207900.] [NT:alternate gene name: yzbD] [LE:130976] [RE:132376] [DI:complement]

ORF Name	NTID	AAID	<u>NT</u> Length!	<u>AA</u> Length	score	probability
AI7503000982_14097011_c3_1411	246	4018	612	203	522	3.6e-50

Description

sp:[LN:ARGJ BACST] [AC:Q07908] [GN:ARGJ] [OR:BACILLUS STEAROTHERMOPHILUS] [EC:2.3.1.35:2.3.1.1] [DE:ACETYLTRANSFERASE, (N-ACETYLGLUTAMATE SYNTHASE) (AGS)] [SP:Q07908] [DB:swissprot] >pir:[LN:I39766] [AC:I39766] [PN:glutamate N-acetyltransferase,] [GN:argJ] [CL:glutamate N-acetyltransferase argJ] [OR:Bacillus stearothermophilus] [EC:2.3.1.35] [DB:pir2] >gp:[GI:g304135] [LN:BACACETYL] [AC:L06036] [PN:ornithine acetyltransferase] [GN:argJ] [OR:Bacillus stearothermophilus] [SR:Bacillus stearothermophilus (strain NCIB 8224) DNA] [DB:genpept-bct1] [EC:2.3.1.35] [DE:Bacillus stearothermophilus ornithine acetyltransferase (argJ) andacetylglutamate kinase (argB) genes, complete cds's, argC gene, 3'end, and argD gene, 5' end.] [NT:also bears acetyl-CoA:L-glutamate] [LE:902] [RE:2134] [DI:direct]

NT AΑ ORF Name NTID AAID score probability LengthLength AI7503000982 14259631 cl 967 4019 1068 355 247 1207 9.3e-123

Description

sp:[LN:QUEA BACSU] [AC:032054] [GN:QUEA] [OR:BACILLUS SUBTILIS] [EC:5.-.-.] [DE: (QUEUOSINE BIOSYNTHESIS PROTEIN QUEA)] [SP:032054] [DB:swissprot] >pir: [LN:A69688] [AC:A69688] [PN:S-adenosylmethionine:tRNA ribosyltransferase-isomerase,:queuosine biosynthesis-related protein queA] [GN:queA] [CL:S-adenosylmethionine:tRNA ribosyltransferase-isomerase] [OR:Bacillus subtilis] [EC: 5.-.-] [DB:pir2] >gp:[GI:e1184021:g2635237] [LN:BSUB0015] [AC:Z99118:AL009126] [PN:S-adenosylmethionine tRNA ribosyltransferase] [GN:queA] [FN:queuosine biosynthesis] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 15 of 21): from 2795131to 3013540.] [SP:032054] [LE:38030] [RE:39058] [DI:complement]

ORF Name	NTID	AAID Length Length score probability
A17503000982_14460932_£1_322	248	4020 177 58
Description gp:[GI:g1022726] [LN:SHU35635] [AC:UI haemolyticus] [SR:Staphylococcus haem [DE:Staphylococcus haemolyticus IS127 [LE:1101] [RE:1922] [DI:complement]	molyticu	
ORF Name	NTID	AAID NT AA score probability
A17503000982_14507827_£1_87	249	4021 150 49
Description NO-HIT		
ORF Name	NTID	AAID NT AA score probability
AI7503000982_14508567_c3_1357	250	4022 276 91 184 2.4e-14
Description		
pir:[LN:E69972] [AC:E69972] [PN:cons [CL:yajC protein] [OR:Bacillus subtil [LN:BSUB0015] [AC:Z99118:AL009126] [OB:genpept-bct1] [DE:Bacillus subtil 2795131to 3013540.] [NT:similar to hy [DI:complement]	lis] [DE GN:yrbF] lis comp	s:pir2] >gp:[GI:e1184019:g2635235] [FN:unknown] [OR:Bacillus subtilis] Plete genome (section 15 of 21): from
ORF Name	NTID	AAID NT AA score probability
Description sp:[LN:GLK_STAXY] [AC:Q56198] [GN:GLE [DE:GLUCOKINASE, (GLUCOSE KINASE)] [S		
[AC:S52352] [PN:glucose kinase] [CL: [OR:Staphylococcus xylosus] [DB:pir2] [PN:glucose kinase] [GN:glkA] [OR:Sta [DE:S.xylosus glucose kinase gene.]	glucose >gp:[Gaphyloco	kinase:glucose kinase homology] [I:g666116] [LN:SXGKG2] [AC:X84332] [Ccus xylosus] [DB:genpept-bct1]
ORF Name	NTID	AAID NT AA score probability
A17503000982_15031535_f2_658	252	4024 [156] [51
Description NO-HIT		
		NT AA
ORF Name	NTID	AAID LengthLength score probability
AI7503000982_15728386_c3_1356	253	4025 1143 380 1542 2.9e-158
<pre>>gp:[GI:e1184020:g2635236] [LN:BSUB00 transglycosylase] [GN:tgt] [FN:queuos</pre>	CION ENZ euine tR [OR:Bac 015] [AC sine bio	YME)] [SP:032053] [DB:swissprot] NA-ribosyltransferase,] [GN:tgt] illus subtilis] [EC:2.4.2.29] [DB:pir2] :Z99118:AL009126] [PN:tRNA-guanine

ORF Name	NTID AAID NT AA score probability
A17503000982_16205035_c1_1036	254 4026 135 44
Description	
NO-HIT	
ORF Name	NTID AAID NT AA score probability
AI7503000982_16610088_c2_1261	255 4027 645 214 405 9.0e-38
Description	
KD PROTEIN IN SERA-FER INTERGENIC REGENCE [AC:E69932] [PN:hypothetical protein >gp:[GI:e1185574:g2634740] [LN:BSUB06 [OR:Bacillus subtilis] [DB:genpept-bc 13 of 21): from 2395261to 2613730.] >gp:[GI:g1146197] [LN:BACSERA] [AC:L6 [DB:genpept-bct2] [DE:Bacillus subtilitypaA,ferredoxin (fer), ypbB, recS, yp (ypcA), ypdA, ypdB, ypdC, spore cortemonophosphatekinase (cmk), ypfD, ypg2	ogenase (glyc), yphE and yphF genes,complete cds.]
ORF Name	NTID AAID NT AA score probability
A17503000982_16828175_c3_1395	256 4028 606 201 1021 4.8e-103
Description	
[OR:Staphylococcus aureus] [DB:genper	F121672] [PN:superoxide dismutase SodA] [GN:sodA] pt-bct2] [DE:Staphylococcus aureus superoxide cds.] [NT:manganese-dependent] [LE:246] [RE:845]
ORF Name	NTID AAID NT AA score probability
A17503000982_16923383_c2_1252	257 4029 339 112 156 3.1e-11
1)] [SP:P54554] [DB:swissprot] reductase homolog yqjQ] [GN:yqjQ] [CR:Bacillus subtilis] [DB:pir2] >gp: [AC:D84432:D82370] [PN:YqjQ] [OR:Bacillus Subtilis] [DB:pir2] [OR:Bacillus Subtilis] [DB:pir2] [OR:Bacillus Subtilis] [DB:pir2] [OR:Bacillus Subtilis] [DB:pir2] [OR:Bacillus Subtilis] [DB:pir2] [OR:Bacillus Subtilis] [DB:pir2] [OR:Bacillus Subtilis] [OB:pir2] [OB:pir2]	

[LE:74971] [RE:75750] [DI:complement]

ORF Name probability

A17503000982 17002217 c2 1254

465

258

4030

154

sp:[LN:FUR2 BACSU] [AC:P54574] [GN:YQKL] [OR:BACILLUS SUBTILIS] [DE:FERRIC UPTAKE REGULATION PROTEIN HOMOLOG 2] [SP:P54574] [DB:swissprot] >pir:[LN:E69967] [AC:E69967] [PN:transcription regulator Fur family homolog yqkL] [GN:yqkL] [CL:ferric uptake regulator] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:d1013328:g1303993] [LN:BACJH642] [AC:D84432:D82370] [PN:YqkL] [OR:Bacillus subtilis] [SR:Bacillus subtilis (strain:JH642(trpC2 PheAl)) DNA] [DB:genpept-bct1] [DE:Bacillus subtilis DNA, 283 Kb region containing skin element.] [LE:269361] [RE:269810] [DI:direct] >qp:[GI:e1185621:g2634787] [LN:BSUB0013] [AC:Z99116:AL009126] [GN:yqkL] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 13 of 21): from 2395261to 2613730.] [NT:similar to transcriptional regulator (Fur family)] [SP:P54574] [LE:53809] [RE:54258] [DI:complement]

ORF Name NTID AAID probability score LengthLength A17503000982 187593 cl 1074 323 3.8e-12

Description

sp:[LN:YPBB_BACSU] [AC:P50728] [GN:YPBB] [OR:BACILLUS SUBTILIS] [DE:HYPOTHETICAL 40.7 KD PROTEIN IN FER-RECQ INTERGENIC REGION] [SP:P50728] [DB:swissprot] >pir:[LN:F69932] [AC:F69932] [PN:hypothetical protein ypbB] [GN:ypbB] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1183748:g2634721] [LN:BSUB0012] [AC:Z99115:AL009126] [GN:ypbB] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 12 of 21): from 2195541to 2409220.] [SP:P50728] [LE:212099] [RE:213157] [DI:complement] >gp:[GI:e1185572:g2634738] [LN:BSUB0013] [AC:Z99116:AL009126] [GN:ypbB] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 13 of 21): from 2395261to 2613730.] [SP:P50728] [LE:12379] [RE:13437] [DI:complement] >gp:[GI:g1146199] [LN:BACSERA] [AC:L47648] [GN:ypbB] [OR:Bacillus subtilis] [DB:genpept-bct2] [DE:Bacillus subtilis phosphoglycerate dehydrogenase (serA), ypaA, ferredoxin (fer), ypbB, recS, ypbD, ypbE, ypbF, ypbG, ypbH, glutamate dehydrogenase (ypcA), ypdA, ypdB, ypdC, spore cortexlytic enzyme (sleB), ypeB, ypfA, ypfB, cytidine monophosphatekinase (cmk), ypfD, ypgA, yphA, yphB, yphC, NAD+ dependentglycerol-3-phosphate dehydrogenase (glyc), yphE and yphF genes,complete cds.] [NT:putative] [LE:3240] [RE:4298] [DI:direct]

NT AA ORF Name NTID AAID probability LengthLength A17503000982_189203_c2_1199 4032 1260 762

Description

sp:[LN:YQEU BACSU] [AC:P54461] [GN:YQEU] [OR:BACILLUS SUBTILIS] [DE:HYPOTHETICAL 28.8 KD PROTEIN IN DNAJ-RPSU INTEREGENIC REGION] [SP:P54461] [DB:swissprot] >pir:[LN:D69952] [AC:D69952] [PN:conserved hypothetical protein yqeU] [GN:yqeU] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:d1013146:g1303811] [LN:BACJH642] [AC:D84432:D82370] [PN:YqeU] [OR:Bacillus subtilis] [SR:Bacillus subtilis (strain:JH642(trpC2 PheA1)) DNA] [DB:genpept-bct1] [DE:Bacillus subtilis DNA, 283 Kb region containing skin element.] [LE:95803] [RE:96573] [DI:direct] >qp:[GI:e1183774:q2634990] [LN:BSUB0014] [AC:Z99117:AL009126] [GN:yqeU] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 14 of 21): from 2599451to 2812870.] [NT:similar to hypothetical proteins] [SP:P54461] [LE:22857] [RE:23627] [DI:complement] >gp:[GI:d1012752:g1890060] [LN:D83717] [AC:D83717] [PN:YqeU] [OR:Bacillus subtilis] [SR:Bacillus subtilis (strain:JH642 (trpC2 pheA1)) DNA] [DB:genpept-bct1] [DE:Bacillus subtilis DNA for DnaJ, YqeT, YqeU, YqeV, YqeW, YqeX, YqeY,complete and partial cds.] [NT:Similar to 26.9 kDa protein (YqqJ) of E. coli] [LE:1334] [RE:2104] [DI:direct]

ORF Name NTID AAID NT AA LengthLength Score probability A17503000982_19540931_c2_1191 Description sp:[LN:CME3_BACSU] [AC:P39695] [GN:COMEC:COME3] [OR:BACILLUS SUBTILIS] [DE:COME OPERON PROTEIN 3] [SP:P39695] [DB:swissprot] >pir:[LN:S39865] [AC:S39865:E69602] [PN:late competence protein 3 (come operon)] [GN:comec] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:g289262] [LN:BACCOME] [AC:L15202] [OR:Bacillus subtilis] [SR:Bacillus subtilis DNA] [DB:genpept-bct2] [DE:Bacillus subtilis come operon encoding ORF1, ORF2, ORF3 andReverse-ORF genes, complete cds.] [LE:2603] [RE:4933] [DI:direct]
NTID AAID LengthLength score probability AI7503000982_19567588_c3_1496 Description sp:[LN:PTGA_BACST] [AC:P42015] [GN:PTSG] [OR:BACILLUS STEAROTHERMOPHILUS] [EC:2.7.1.69] [DE:COMPONENT), (EII-GLC / EIII-GLC) (FRAGMENT)] [SP:P42015] [DB:swissprot] >gp:[GI:g529001] [LN:BSU12340] [AC:U12340] [PN:PTS glucose-specific permease] [GN:ptsG'] [OR:Bacillus stearothermophilus] [DB:genpept-bct1] [DE:Bacillus stearothermophilus XL-65-6 phosphoenolpyruvate-dependentphosphotransferase system glucose-specific permease (ptsG') gene, partial cds, HPr (ptsH), enzyme I (ptsI), and PtsT (ptsT) genes, complete cds, and wall associated protein precursor (wapA') gene, complete cds.] [NT:thermophilic, cytoplasmic protein] [LE:<1] [RE:976] [DI:direct]
ORF Name NTID AAID NT AA LengthLength LengthLength LengthLength LengthLength Description Sp: [LN:YPPQ_BACSU] [AC:P54155] [GN:YPPQ] [OR:BACILLUS SUBTILIS] [DE:HYPOTHETICAL 16.6 KD PROTEIN IN ILVA 3'REGION] [SP:P54155] [DB:swissprot] >pir: [LN:F69940] [AC:F69940] [PN:transcription regulator PilB family homolog yppQ] [GN:yppQ] [CL:hypthetical protein YCL033c] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:g1256654] [LN:BACYACA] [AC:L77246] [GN:yppQ] [OR:Bacillus subtilis] [DB:genpept-bctl] [DE:Bacillus subtilis (YAC10-9 clone) DNA region between the serA andkdg loci.] [NT:54.8% identity with Neisseria gonorrhoeae] [LE:33923] [RE:34354] [DI:direct] >gp:[GI:e1183615:g2634588] [LN:BSUB0012] [AC:Z99115:AL009126] [GN:yppQ] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bctl] [DE:Bacillus subtilis complete genome (section 12 of 21): from 2195541to 2409220.] [NT:similar to transcriptional regulator (PilB family)] [SP:P54155] [LE:90798] [RE:91229] [DI:complement]
ORF Name NTID AAID NT AA LengthLength score probability A17503000982_1960017_f3_809 264 4036 135 44

ORF Name	NTID AAID NT AA score probability
A17503000982_19804838_f2_385	7
KD PROTEIN IN BCSA-DEGR INTERGENIC [AC:C69934] [PN:conserved hypothet fulgidus conserved hypothetical pro >gp:[GI:g1256625] [LN:BACYACA] [AC:[DB:genpept-bct1] [DE:Bacillus subt andkdg loci.] [NT:putative] [LE:110 >gp:[GI:e1183645:g2634618] [LN:BSUB [OR:Bacillus subtilis] [DB:genpept-	0012] [AC:Z99115:AL009126] [GN:ypdP] [FN:unknown] bctl] [DE:Bacillus subtilis complete genome (section [NT:similar to hypothetical proteins] [SP:P54163]
ORF Name	$rac{ ext{NTID}}{ ext{AAID}} = rac{ ext{NT}}{ ext{Length}} rac{ ext{AA}}{ ext{score}} = rac{ ext{probability}}{ ext{probability}}$
KD PROTEIN IN CCCA-SODA INTERGENIC I [AC:H69953] [PN:conserved hypothet. [DB:pir2] >gp:[GI:d1013170:g1303835] [OR:Bacillus subtilis] [SR:Bacillus [DB:genpept-bct1] [DE:Bacillus subtilis] [LE:120338] [RE:120988] [DI:direct] [AC:Z99116:AL009126] [GN:yqfN] [FN:u] [DE:Bacillus subtilis complete genor	YQFN] [OR:BACILLUS SUBTILIS] [DE:HYPOTHETICAL 23.7 REGION] [SP:P54471] [DB:swissprot] >pir:[LN:H69953] ical protein yqfN] [GN:yqfN] [OR:Bacillus subtilis] [LN:BACJH642] [AC:D84432:D82370] [PN:YqfN] subtilis (strain:JH642(trpC2 PheA1)) DNA] ilis DNA, 283 Kb region containing skin element.] >gp:[GI:e1185785:g2634951] [LN:BSUB0013] unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] me (section 13 of 21): from 2395261to 2613730.] s] [SP:P54471] [LE:202632] [RE:203282]
ORF Name	NTID AAID Longth cont probability
AI7503000982_20035967_c1_1082	Length Length 5501e 510535111ty
[GN:gpsA] [CL:glycerol-3-phosphate [DB:pir2] >gp:[GI:e1183728:g2634701] [PN:NAD(P)H-dependent glycerol-3-phosphate] [OR:Bacillus subtilis] subtilis complete genome (section 12 [LE:192848] [RE:193885] [DI:compleme [PN:NAD+ dependent glycerol-3-phosphate] [DB:genpept-bct2] [EC:1.1.1.94] [DE:(serA), ypaA,ferredoxin (fer), ypbB, dehydrogenase (ypcA), ypdA, ypdB, ypypfB, cytidine monophosphatekinase	ycerol-3-phosphate dehydrogenase (NAD+), gpsA] dehydrogenase] [OR:Bacillus subtilis] [EC:1.1.1.8]] [LN:BSUB0012] [AC:Z99115:AL009126] osphate] [GN:gpsA] [FN:synthesis of the sn-glycerol [DB:genpept-bct1] [EC:1.1.1.94] [DE:Bacillus 2 of 21): from 2195541to 2409220.] [SP:P46919] ent] >gp:[GI:g1146220] [LN:BACSERA] [AC:L47648] hate] [GN:glyC] [OR:Bacillus subtilis] :Bacillus subtilis phosphoglycerate dehydrogenase , recS, ypbD, ypbE, ypbF, ypbG, ypbH,glutamate pdC, spore cortexlytic enzyme (sleB), ypeB, ypfA, (cmk), ypfD, ypgA, yphA, yphB, yphC, NAD+
[LE:22512] [RE:23549] [DI:direct]	rogenase (glyc), yphE and yphF genes,complete cds.]

NT AA Length Length score AAID ORF Name NTID probability A17503000982 2038325 £2 381 4041 180 269 59 Description NO-HIT NTAA ORF Name NTID AAID score probability LengthLength AI7503000982 20485712 c3 1498 270 4042 **1573** Description NO-HIT NT AAORF Name NTID AAID score probability LengthLength AI7503000982 20503437 c3 1456 271 4043 123 40 Description NO-HIT NT AΑ ORF Name NTID AAID score probability LengthLength AI7503000982_20507937_c2_1280 272 4044 1317 438 1823 4.9e-188 Description sp:[LN:SYN_BACSU] [AC:P39772] [GN:ASNS] [OR:BACILLUS SUBTILIS] [EC:6.1.1.22] [DE:(ASNRS)] [SP:P39772] [DB:swissprot] >pir:[LN:B69591] [AC:B69591:I40523] [PN:asparagine--tRNA ligase, asnS:asparaginyl-tRNA synthetase asnS] [GN:asnS] [CL:lysine--tRNA ligase] [OR:Bacillus subtilis] [EC:6.1.1.22] [DB:pir2] >gp:[GI:g1146247] [LN:BACYPIA] [AC:L47709] [PN:asparaqinyl-tRNA synthetase] [GN:asnS] [OR:Bacillus subtilis] [DB:genpept-bct1] [EC:6.1.1.22] [DE:Bacillus subtilis (clone YAC15-6B) ypiABF genes, qcrABC genes, ypjABCDEFGHI genes, birA gene, panBCD genes, dinG gene, ypmB gene, aspB gene, asnS gene, dnaD gene, nth gene and ypoC gene, completecds's.] [NT:41.1% of identity to the Escherichia coli] [LE:20449] [RE:21741] [DI:direct] >gp:[GI:e1183681:g2634654] [LN:BSUB0012] [AC:Z99115:AL009126]

[PN:asparaginyl-tRNA synthetase] [GN:asnS] [OR:Bacillus subtilis] [DB:genpept-bct1] [EC:6.1.1.22] [DE:Bacillus subtilis complete genome (section 12 of 21): from 2195541to

2409220.] [SP:P39772] [LE:149926] [RE:151218] [DI:complement]

ORF Name probability Length Length AI7503000982 20517135 c3 1457 315 104 sp:[LN:DBH BACST] [AC:P02346:P08822] [GN:HBS:HBSU] [OR:BACILLUS STEAROTHERMOPHILUS: BACILLUS CALDOLYTICUS: BACILLUS CALDOTENAX] [DE:DNA-BINDING PROTEIN II (HB) (HU)] [SP:P02346:P08822] [DB:swissprot] >pir:[LN:DNBS2F] [AC:JC1205:A02690:JC2509] [PN:DNA-binding protein HU:DNA-binding protein II] [GN:hup] [CL:bacterial DNA-binding protein] [OR:Bacillus stearothermophilus] [DB:pir1] >pir:[LN:JC1207] [AC:JC1207] [PN:DNA-binding protein HU] [GN:hup] [CL:bacterial DNA-binding protein] [OR:Bacillus caldolyticus] [DB:pir2] >pir:[LN:JC1206] [AC:JC1206] [PN:DNA-binding protein HU] [GN:hup] [CL:bacterial DNA-binding protein] [OR:Bacillus caldotenax] [DB:pir2] >gp:[GI:d1007851:g1065992] [LN:BACDBPHU] [AC:D38080] [PN:DNA binding protein HU] [OR:Bacillus stearothermophilus] [SR:Bacillus stearothermophilus (strain:1503) DNA] [DB:genpept-bct1] [DE:Bacillus stearothermophilus gene for DNA binding protein HU, complete cds.] [LE:13] [RE:285] [DI:direct] >gp:[GI:g143065] [LN:BACHUB1] [AC:M73500] [PN:hubst] [GN:hubst] [OR:Bacillus stearothermophilus] [SR:Bacillus stearothermophilus DNA] [DB:genpept-bct1] [DE:B.stearothermophilus hubst gene, complete cds.] [LE:1] [RE:>270] [DI:direct] >qp:[GI:q143067] [LN:BACHUB2] [AC:M73501] [PN:hubcalx] [GN:hubcalx] [OR:Bacillus caldotenax] [SR:Bacillus caldotenax DNA] [DB:genpept-bct1] [DE:B.caldotenax hubcalx gene, 5' end.] [LE:1] [RE:>270] [DI:direct] >gp:[GI:g143069] [LN:BACHUB3] [AC:M73502] [PN:hubcald] [GN:hubcald] [OR:Bacillus caldolyticus] [SR:Bacillus caldolyticus DNA] [DB:qenpept-bct1] [DE:B.caldolyticus hubcald gene, 5' end.] [LE:1] [RE:>270] [DI:direct] AA ORF Name NTID AAID probability Length Length AI7503000982 20587536 f1 4 Description sp:[LN:XPAC BACSU] [AC:P37467] [GN:XPAC] [OR:BACILLUS SUBTILIS] [DE:XPAC PROTEIN] [SP:P37467] [DB:swissprot] >pir:[LN:S27526] [AC:S27526:S66055:C69734] [PN:5-bromo-4-chloroindolyl phosphate hydrolysis protein xpaC:xpaC protein] [GN:xpaC] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:d1005803:g467415] [LN:BAC180K] [AC:D26185] [PN:hydrolysis of 5-bromo-4-chloroindolyl phosphate] [GN:xpaC] [OR:Bacillus subtilis] [SR:Bacillus subtilis (sub species:Marburg, strain:168) DNA] [DB:genpept-bct1] [DE:B. subtilis DNA, 180 kilobase region of replication origin.] [LE:99445] [RE:100059] [DI:direct] >gp:[GI:g143830] [LN:BACXPAC] [AC:M96156] [GN:xpaC] [OR:Bacillus subtilis] [SR:Bacillus subtilis (strain W168) DNA] [DB:genpept-bct1] [DE:Bacillus subtilis ORF1 and xpaC gene, complete cds's; ssrRNA gene,3' end; ORF3C 5' end.] [LE:532] [RE:1146] [DI:direct] >gp:[GI:e1181958:g2632292] [LN:BSUB0001] [AC:Z99104:AL009126] [GN:xpaC] [FN:hydrolysis of 5-bromo 4-chloroindolyl phosphate] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 1 of 21): from 1 to213080.] [SP:P37467] [LE:35843] [RE:36457] [DI:direct] NT ORF Name NTID LengthLength 275 Description

AI7503000982 20594688 c2 1331

sp:[LN:LPLA BACSU] [AC:P37966] [GN:LPLA] [OR:BACILLUS SUBTILIS] [DE:LIPOPROTEIN LPLA PRECURSOR] [SP:P37966] [DB:swissprot] >pir:[LN:I39876] [AC:I39876:H69652] [PN:lipoprotein lplA:lysis protein lplA] [GN:lplA] [CL:Bacillus subtilis lipoprotein lplA] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1182689:g2633023] [LN:BSUB0004] [AC:Z99107:AL009126] [PN:lipoprotein] [GN:lplA] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 4 of 21): from 600701 to813890.] [SP:P37966] [LE:178337] [RE:179845] [DI:direct] >qp:[GI:q431272] [LN:BACLPLA] [AC:L03376] [PN:lysis protein] [GN:lplA] [OR:Bacillus subtilis] [SR:Bacillus subtilis (strain Marburg 168) DNA] [DB:genpept-bct2] [DE:Bacillus subtilis lysis protein (lplA) gene, complete cds.] [LE:482] [RE:1990] [DI:direct]

ORF Name	NTID	AAID	NT AA score probability
A17503000982_20734677_c1_1135	276	4048	774 257 297 2.5e-26
Description gp:[GI:g4981613] [LN:AE001767] [AC:A DeoR family] [GN:TM1069] [OR:Thermot maritima section 79 of 136 of the co identity: 55.02;] [LE:1578] [RE:2336	oga mar omplete	itima] genome	[DB:genpept-bct2] [DE:Thermotoga .] [NT:similar to GB:AL009126 percent
ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000982_20791068_c3_1370	277	4049	273 90 71 0.041
Description gp:[GI:g294060] [LN:PAPMPL146A] [AC: [OR:Papaver somniferum] [SR:Papaver [DB:genpept-pln1] [DE:Papaver somnif cds.] [LE:963:1290] [RE:1167:1564] [somnife: erum ma	rum (st jor lat	train UNL186) (library: EMBL) DNA] tex protein (MLP146) gene, complete
ORF Name AI7503000982_20876263_f1_159 Description NO-HIT	NTID 278	<u>AAID</u> 4050	NT AA Score probability LengthLength 56
ORF Name	NTID	AAID	NT AA score probability
Description sp:[LN:AROC_STAAU] [AC:Q59803] [GN:A [DE:PHOSPHOLYASE)] [SP:Q59803] [DB:s [AC:U31979] [PN:chorismate synthase] [DB:genpept-bct1] [EC:4.6.1.4] [DE:S nucleosidediphosphate kinase (ndk) g and geranylgeranyl pyrophosphate syn [NT:5-enolpyruvylshikimate 3-phospha	wissprog [GN:arc staphylogenes, co thetase	t] >gp oC] [OH coccus omplete homolo	:[GI:g987498] [LN:SAU31979] R:Staphylococcus aureus] aureus chorismate synthase (aroC) and e cds, dehydroauinatesynthase (aroB) og(gerCC) genes, partial cds.]
ORF Name A17503000982_21501550_c1_1127	NTID 280	<u>AAID</u> 4052	NT AA LengthLength score probability [1116] 371 [999] [1.0e-100]
Description gp:[GI:g3688811] [LN:AF084104] [AC:A protein] [GN:malK] [OR:Bacillus firm (acsA) gene, partial cds; SspA (sspA ATP-binding protein(malK), leucine-r (lrpR), hypothetical proteins, ABC tr NatB (natB), and hypothetical protei partial cds.] [NT:MalK; Orf4; simila [RE:3490] [DI:complement]	nus] [DB A),hypoth ich prof ansporte n genes	genper hetical tein to er ATP- , compl	pt-bct2] [DE:Bacillus firmus AcsA l protein, maltose transportor ranscriptional regulator -binding protein (natC),NatA (natA), letecds; and SpoIIIJ (spoIIIJ) gene,
ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000982_21526562_c2_1206	281	4053	138 45
Description NO-HIT			

ORF Name	NTID	AAID	NT AA Length Length	probability
A17503000982_21531627_c3_1483	282	4054	132 43	
Description		L		
NO-HIT				
ORF Name	NTID	AAID	NT AA LengthLength	probability
AI7503000982_21537962_c1_1093	283	4055	612 203 357 1	1e-32
Description				
sp:[LN:YPJA_BACSU] [AC:P54392] [GN:YIKD PROTEIN IN QCRC-DAPB INTERGENIC RI[AC:A69937] [PN:hypothetical protein >gp:[GI:g1146230] [LN:BACYPIA] [AC:L4subtilis] [DB:genpept-bct1] [DE:Bacilgenes, ypjABCDEFGHI genes, birA gene, gene, dnaD gene, nth gene and ypoC gene; dnaD gene, nth gene and ypoC gene; [GI:e118369] [GN:ypjA] [FN:unknown] [OR:Bacillus stomplete genome (section 12 of 21): fene [RE:166665] [DI:complement]	EGION] I ypjA] I ypjA] Ilus suk panBCD Ene, com 98:g2634	[SP:P5- [GN:yp: ctilis genes mpleted [671]	A392] [DB:swissprot] pjA] [OR:Bacillus s jA] [FN:hypothetica] (clone YAC15-6B) yp , dinG gene, ypmB ge cds's.] [NT:putative [LN:BSUB0012] [AC:ZS :genpept-bct1] [DE:E	>pir:[LN:A69937] subtilis] [DB:pir2] L] [OR:Bacillus DIABF genes, qcrABC Ene,aspB gene, asnS E] [LE:5002] D9115:AL009126] Bacillus subtilis
ORF Name	NTID	AAID	NT AA LengthLength	probability
A17503000982_21568762_c1_1013	284	4056	1005 334 992 5	.6e-100
Description			J	·
<pre>gp:[GI:e1356351:g3947511] [LN:BMAJ482 megaterium] [DB:genpept-bct1] [DE:Bac region, 3'flanking region.] [LE:3056]</pre>	cillus n	negate	rium DSM319 spoIV op	
ORF Name	NTID	AAID	NT AA LengthLength	probability
A17503000982_21600325_f1_228	285	4057	141 46	
Description NO-HIT				
ORF Name	NTID	AAID	NT <u>AA</u> LengthLength	probability
A17503000982_21640636_f1_136	286	4058	276 91 366 1	.2e-33
Description		-		
pir: [LN:D69621] [AC:D69621] [PN:ferr 2[4Fe-4S]:ferredoxin 2[4Fe-4S] homolo >gp: [GI:e1183749:g2634722] [LN:BSUB00 [OR:Bacillus subtilis] [DB:genpept-bc 12 of 21): from 2195541to 2409220.] [LE:213423] [RE:213671] [DI:direct] > [AC:Z99116:AL009126] [PN:ferredoxin] [DE:Bacillus subtilis complete genome [NT:alternate gene name: ypbA] [SP:P5 >gp: [GI:g1146198] [LN:BACSERA] [AC:L4 subtilis] [DB:genpept-bct2] [DE:Bacil (serA), ypaA, ferredoxin (fer), ypbB, dehydrogenase (ypcA), ypdA, ypdB, ypd ypfB, cytidine monophosphatekinase (cdependentglycerol-3-phosphate dehydrogense [NT:ypbA; similar to B.stearothermoph [DI:complement]	pgy] [OR D12] [AC ct1] [DE [NT:alte gp:[GI: [GN:fer composed [GN:fer areas] [GN:fer composed [GN:fer areas] [GN	E:Bacil E:Z9911 E:Bacil Ernate e11855 [OR: on 13 [LE:137 Exilis TPDD, ye cort ofD, ye (glyc)	Llus subtilis] [DB:p 15:AL009126] [PN:fer Llus subtilis comple gene name: ypbA] [S 173:g2634739] [LN:BS 18acillus subtilis] of 21): from 239526 1703] [RE:13951] [DI: 17redoxin] [GN:fer] [17phosphoglycerate de 17pbE, ypbF, ypbG, yp 18exlytic enzyme (sle 19gA, yphA, yphB, yph 19gA, yphE and yphF gen	redoxin] [GN:fer] redoxin] [GN:fer] re genome (section P:P50727] UB0013] [DB:genpept-bct1] 1to 2613730.] direct] OR:Bacillus hydrogenase bH,glutamate B), ypeB, ypfA, C, NAD+ es,complete cds.]

			NT A	A A	
ORF Name	NTID	AAID	LengthLen		probability
AI7503000982_21667676_c1_975	287	4059	888 29	5 590	2.2e-57
Description		<u></u>			
pir:[LN:C69981] [AC:C69981] [PN:con [OR:Bacillus subtilis] [DB:pir2] >gp [AC:Z99118:AL009126] [GN:yrvM] [FN:u [DE:Bacillus subtilis complete genom [NT:similar to hypothetical proteins	:[GI:e1: nknown] e (sect:	184002 OR:B ion 15	:g2635218] acillus su of 21): f	[LN:BSUB ubtilis] [from 27951	0015] DB:genpept-bct1] 31to 3013540.]
ORF Name	NTID	AAID	<u>NT A</u> LengthLen	<u>A</u> gth	probability
AI7503000982_21674067_f1_8	288	4060	198 65		
Description					
NO-HIT					
ORF Name	NTID	AAID	NT A	A gth score	probability
A17503000982_21678187_c3_1486	289	4061	240 79		
Description		•	<u> </u>		
NO-HIT					
ORF Name	NTID	AAID	NT A LengthLen	~ caara	probability
AI7503000982_21730443_c1_1055	290	4062	1332 443	904	1.2e-90
Description					
sp:[LN:ODB2_BACSU] [AC:P37942] [GN:B3 [EC:2.3.1] [DE:CHAIN TRANSACYLASE] [AC:S32488:E69593] [PN:dihydrolipoard dehydrogenase complex (bfmBB):branched component:dihydrolipoyl acyltransfer acetyltransferase: lipoyl/biotin-bind [DB:pir2] >gp:[GI:g142613] [LN:BACBR alpha-keto acid dehydrogenase E2] [OB:genpept-bct1] [DE:Bacillus subtited dehydrogenase E1-alpha, branched chain and branched chain alpha-keto acid del [DI:direct] >gp:[GI:d1013279:g1303944]] [SP:P3 mide S-a ed-chair ase] [GN ding hom ANCH] [A R:Bacil] lis bran n alpha-	37942] acyltra alpha 1:bfmBl mology AC:M973 lus sul ached a -keto a	[DB:swiss ansferase, a-oxo acid B] [CL:di] [OR:Baci 391:M96937 btilis] [Schain alphacid dehyd 2, complet	sprot] >pi , alpha-o l dehydrog hydrolipo llus subt] [PN:bra R:Bacillu a-keto ac lrogenase e cds.] [r:[LN:S32488] xo acid enase complex E2 amide ilis] [EC:2.3.1] nched chain s subtilis DNA] id E1-beta, LE:2228] [RE:3502]
[OR Bacillus subtilis] [SR Bacillus of	zuhtilie	e (etra	in .TW642/	trnC2 Dha	אוו און א

[DB:genpept-bct1] [DE:Bacillus subtilis DNA, 283 Kb region containing skin element.]

[AC:Z99116:AL009126] [PN:branched-chain alpha-keto acid dehydrogenase E2] [GN:bfmBB] [OR:Bacillus subtilis] [DB:genpept-bct1] [EC:2.3.1.-] [DE:Bacillus subtilis complete genome (section 13 of 21): from 2395261to 2613730.] [NT:alternate gene name: bfmB2]

[LE:221574] [RE:222848] [DI:direct] >gp:[GI:e1185671:g2634837] [LN:BSUB0013]

[SP:P37942] [LE:100771] [RE:102045] [DI:complement]

	NTID	AAID	NT LengthL	<u>AA</u> .ength ^S	core	probabilit	Y
A17503000982_21756562_c2_1289	291	4063	696 [2	231	162	5.1e-12	
Description		L	JL L				
pir: [LN:G69828] [AC:G69828] [PN:cal [OR:Bacillus subtilis] [DB:pir2] >gp [AC:Z99109:AL009126] [GN:yheG] [FN:u [DE:Bacillus subtilis complete genom [NT:similar to calcium-binding prote >gp:[GI:e325187:g2226164] [LN:BSY140 [OR:Bacillus subtilis] [DB:genpept-b75 degrees: sspB upstreamof glyB.] [LE:11988] [RE:12608] [DI:direct]	o:[GI:e1] unknown] ne (sect. ein] [LE 080] [AC oct1] [D	182974 [OR:Baion 6 (:49122] :Y1408(E:Bacil	g263330 acillus of 21): [RE:49] [PN:h	08] [LN subtil from 90742] [aypothe btilis	:BSUB is] [99501 DI:co tical chrom	0006] DB:genpept-k to1209940.] mplement] protein] [Goognath of the content of t	oct1] N:yheG] region
ORF Name	NTID	AAID	<u>NT</u> LengthL	ength	core	probabilit	Υ
A17503000982_21759427_c3_1407 Description	292	4064	1350 4	149	1396	3.7e-143	
sp:[LN:GCS1_BACSU] [AC:P54376] [GN:Y [DE:PROTEIN)] [SP:P54376] [DB:swissp dehydrogenase homolog yqhJ] [GN:yqhJ >gp:[GI:d1013226:g1303891] [LN:BACJH subtilis] [SR:Bacillus subtilis (str [DE:Bacillus subtilis DNA, 283 Kb re [RE:172736] [DI:direct] >gp:[GI:e118 [GN:yqhJ] [FN:unknown] [OR:Bacillus complete genome (section 13 of 21): dehydrogenase] [SP:P54376] [LE:15088	orot] >p: [OR:1] [642] [AG (ain:JH64) (gion con (5724:g26) (subtilis (from 239)	ir:[LN: Bacillu C:D8443 42(trp0 ntainir 534890] B] [DB:	:A69959] IS SUBTI B2:D8237 C2 PheA1 Ing Skin [LN:BS Egenpept D 261373	[AC:A lis] [: 0] [PN .)) DNA elemen :UB0013 bct1]	69959 DB:pi :YqhJ] [DB t.] []] [AC [DE:] T:sim] [PN:glyci r2]] [OR:Bacill :genpept-bct LE:171390] :Z99116:AL00 Bacillus sub	us 1] 9126] tilis
ORF Name	NTID	AAID	NT	AA S	core	1 1 1 1 1 1 1 1	
			LengthL	ength 🖰	COIC	probabilit	Y
A17503000982_21914067_c3_1512	293	4065	LengthL	ength		probabilit 3.1e-51	<u>Y</u>
AT7503000982_21914067_c3_1512 Description gp:[GI:g2182835] [LN:LLU81166] [AC:U [OR:Lactococcus lactis subsp. cremor cremoris MG1363 histidine kinase (ll [DI:direct]	[81166] [DB:	4065 [PN:his	1416 4 stidine ot-bct2]	kinase	Llkii	nA] [GN:llki	nA]
Description gp:[GI:g2182835] [LN:LLU81166] [AC:U [OR:Lactococcus lactis subsp. cremor cremoris MG1363 histidine kinase (11	[81166] [DB:	4065 [PN:his:genper	1416 4 stidine ot-bct2]	kinase [DE:Lads.] [1	Llkii	nA] [GN:llki	nA] subsp.
Description gp:[GI:g2182835] [LN:LLU81166] [AC:U [OR:Lactococcus lactis subsp. cremor cremoris MG1363 histidine kinase (ll [DI:direct]	B1166] ris] [DB: kinA)ger	4065 [PN:his:genper	stidine st-bct2] aplete c	kinase [DE:Lads.] [I	Llkinactoco LE:1]	nA] [GN:llki occus lactis [RE:1473]	nA] subsp.
Description gp:[GI:g2182835] [LN:LLU81166] [AC:U [OR:Lactococcus lactis subsp. cremor cremoris MG1363 histidine kinase (ll [DI:direct] ORF Name	MED NOTED	4065 [PN:his:genper] ne, con	stidine st-bct2] aplete c NT Length Le	kinase [DE:Ladds.] [1 AA ength 64 3	Llkinactocc LE:1]	A.le-51 (GN:llki) (CCCUS lactis) (RE:1473) probability (P.9e-32)	nA] subsp.

[OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 13 of 21): from 2395261to 2613730.] [SP:P25954] [LE:162000] [RE:162971] [DI:complement]

ORF Name	NTID AAID NT Lengtl	AA hLength score	probability
AI7503000982_21962762_c1_1139	295 4067 732	243 615 5	.0e-60
Description		/	
<pre>gp:[GI:g143267] [LN:BACODHAB] [AC:M2 3G18) DNA] [DB:genpept-bct1] [DE:B.s end, anddihydrolipoamide transsuccin [NT:2-oxoglutarate dehydrogenase (od</pre>	ubtilis 2-oxoglutar vlase (odhB) gene,	ate dehydrogen complete cds.]	ase (odhA) gene 3'
ORF Name	NTID AAID NT Length	AA hLength	probability
A17503000982_22031307_£3_906	296 4068 150	49	
Description		, <u> </u>	
NO-HIT			
ORF Name	NTID AAID NT Length	<u>AA</u> nLength score	probability
A17503000982_2227312_c1_974	297 4069 1782	593 2019 8	.4e-209
Description			
[DE:(ASPRS)] [SP:032038] [DB:swisspr [PN:aspartatetRNA ligase, aspS:asp ligase] [OR:Bacillus subtilis] [EC:6 [LN:BSUB0015] [AC:Z99118:AL009126] [subtilis] [DB:genpept-bct1] [EC:6.1. (section 15 of 21): from 2795131to 3 [DI:complement]	rtyl-tRNA syntheta 1.1.12] [DB:pir2] PN:aspartyl-tRNA sy 12] [DE:Bacillus	se] [GN:aspS] >gp:[GI:e11840 nthetase] [GN: subtilis compl	03:g2635219] aspS] [OR:Bacillus ete genome
ORF Name	NTID AAID NT Length	AA nLength	probability
A17503000982_22459462_c2_1275	298 4070 138	45	
Description			
NO-HIT			
ORF Name	NTID AAID NT Length	AA nLength	probability
A17503000982_22661088_c3_1363	299 4071 150	49	
Description			
NO-HIT			
ORF Name	NTID AAID NT Length	AA Length score	probability
A17503000982_23392_f2_366	300 4072 147	48	
Description			
NO-HIT			

ORF Name	NTID	<u>AAID</u>	NT Length	<u>AA</u> Length	score	probability
AI7503000982_23445266_c1_1120	301	4073		188		3.3e-56
Description	L	<u> </u>			J ()	
sp:[LN:PMSR_BACSU] [AC:P54154] [GN:Y [SP:P54154] [DB:swissprot] >pir:[LN: sulfoxide reductase homolog yppP] [Greductase] [OR:Bacillus subtilis] [D [PN:DNA-binding protein] [GN:yppP] [DE:Bacillus subtilis (YAC10-9 clone [NT:42.4% identity with the Lycopers >gp:[GI:e1183616:g2634589] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-b 12 of 21): from 2195541to 2409220.] reductase] [SP:P54154] [LE:91230] [R	E69940] N:yppP B:pir2] OR:Baci) DNA ro icon eso 012] [Ac ct1] [DI	[AC:E] [CL: >gp:[llus s egion culent C:Z991 E:Baci ilar t	69940] peptide GI:g125 ubtilis between um] [LE 15:AL00 llus su o pepti	[PN:permething of the second o	peptide conine [LN:BA genper gerA ar B] [RE: [GN:YI] compl	e methionine sulfoxide ACYACA] [AC:L77246] ot-bct1] adkdg loci.] 33922] [DI:direct] opp] [FN:unknown] ete genome (section
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000982_23445762_c3_1397	302	4074	144	47]	
Description					_	
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
AI7503000982_23468938_£1_258	303	4075	168	55]	
Description					-	
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000982_23470327_c2_1225	304	4076	1461	486	446	4.1e-42
Description pir:[LN:S52351] [AC:S52351] [PN:hype [DB:pir2] >gp:[GI:g666115] [LN:SXGKG] [DB:genpept-bct1] [DE:S.xylosus gluckinase] [LE:<1] [RE:406] [DI:direct]	2] [AC:	(84332)] [GN:u	gl] [C	R:Stap	hylococcus xylosus]
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
AI7503000982_23470452_c2_1183	305	4077	441	146	214	1.6e-17
Description pir:[LN:F71860] [AC:F71860] [PN:biographic carboxyl carrier protein: lipoyl/biographic carboxyl carrier protein: lipoyl/biographic carboxyl carrier protein: lipoyl/biographic carboxyl carrier protein: lipoyl/biographic carboxyl carrier protein: lipoyl carboxyl	tin-bind rain J99 [PN:BI0 pept-bct	ding horized to the control of the c	omology OB:pir2 ARBOXYL E:Helic] [OR:] >gp: CARRI obacte	Helico [GI:g4 ER PRO r pylo	bacter pylori] 155592] TEIN] [GN:accB] ri, strain J99
ORF Name	NTID	AAID	<u>NT</u> Lengthl	<u>AA</u> Length	<u>score</u>	probability
AI7503000982_23476676_c2_1204	306	4078	369	122	227	6.6e-19
Description gp:[GI:g902055] [LN:BSU29177] [AC:U2: [OR:Bacillus subtilis] [DB:genpept-bottomical cds, diacylglycerolkinase (dd.) (cdd) gene,partial cds.] [LE:2793] [Formula comment of the comment of t	ct1] [DE gk) gene	E:Bacil	llus sul plete c	btilis	PhoH	(phoH) gene,

ORF Name	NTID AAID LengthLength score probability	
AI7503000982_23595137_c2_1185	307 4079 558 185 522 3.6e-50	
Description sp:[LN:YQEG_BACSU] [AC:P54452] [GN:Y KD PROTEIN IN NUCB-AROD INTERGENIC R [AC:C69951] [PN:conserved hypotheti [DB:pir2] >gp:[GI:d1013122:g1303787] [OR:Bacillus subtilis] [SR:Bacillus [DB:genpept-bct1] [DE:Bacillus subti [LE:72502] [RE:73020] [DI:direct] >g [AC:Z99117:AL009126] [GN:yqeG] [FN:U [DE:Bacillus subtilis complete genom [NT:similar to hypothetical proteins	[SP:P54452] [LE:46410] [RE:46928] [DI:comple	9951] tilis] t.] t1]
ORF Name A17503000982_23597252_f2_557 Description NO-HIT	NTID AAID LengthLength score probability 308 4080 168 55	
ORF Name A17503000982_23620205_f2_640 Description NO-HIT	NTID AAID NT AA score probability 309 4081 138 45	
<pre>[EC:3.2.1.20] [DB:pir2] >gp:[GI:g474 [PN:alpha-D-1,4-glucosidase] [GN:mal</pre>	ore homology] [OR:Staphylococcus xylosus]	
ORF Name A17503000982_23626383_t3_761 Description NO-HIT	NTID AAID NT AA score probability LengthLength 311 4083 132 43	
ORF Name A17503000982_23626425_c1_1042 Description pir: [LN:S72490] [AC:S72490:I39765] [GN:argC] [CL:N-acetyl-gamma-glutam stearothermophilus] [EC:1_2_1_38] [D		se,]

ODE Name	NT AA				
ORF Name	NTID AAID LengthLength score probability				
A17503000982_23642942_c1_1012	313 4085 696 231 142 3.6e-07				
[OR:Plasmodium falciparum] [DB:pir2] [AC:AE001410:AE001362] [PN:hypothet falciparum] [SR:malaria parasite P. falciparum chromosome 2, section 47	pothetical protein PFB0680w] [GN:PFB0680w]] >gp:[GI:g3845248] [LN:AE001410] ical protein] [GN:PFB0680w] [OR:Plasmodium falciparum] [DB:genpept-inv2] [DE:Plasmodium of 73 of thecomplete sequence.] [NT:predicted by E:10567:12528:12807] [DI:directJoin]				
ORF Name	NTID AAID <u>NT AA</u> score probability				
A17503000982_23703452_c1_1037	314 4086 1509 502 1808 1.9e-186				
Description sp:[LN:GCS2_BACSU] [AC:P54377] [GN:YQHK] [OR:BACILLUS SUBTILIS] [EC:1.4.4.2] [DE:PROTEIN)] [SP:P54377] [DB:swissprot] >pir:[LN:B69959] [AC:B69959] [PN:glycine dehydrogenase homolog yqhK] [GN:yqhK] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:d1013227:g1303892] [LN:BACJH642] [AC:D84432:D82370] [PN:YqhK] [OR:Bacillus subtilis] [SR:Bacillus subtilis (strain:JH642(trpC2 PheA1)) DNA] [DB:genpept-bct1] [DE:Bacillus subtilis DNA, 283 Kb region containing skin element.] [LE:172729] [RE:174195] [DI:direct] >gp:[GI:e1185723:g2634889] [LN:BSUB0013] [AC:Z99116:AL009126] [GN:yqhK] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 13 of 21): from 2395261to 2613730.] [NT:similar to glycine dehydrogenase] [SP:P54377] [LE:149426] [RE:150892] [DI:complement]					
ORF Name	NTID AAID NT AA score probability				
A17503000982_23707890_c2_1322 Description	315 4087 240 79 123 6.9e-08				
<pre>subtilis] [DB:pir2] >gp:[GI:e1185439 [GN:yozE] [FN:unknown] [OR:Bacillus</pre>	pothetical protein yozE] [GN:yozE] [OR:Bacillus 9:g2634360] [LN:BSUB0011] [AC:Z99114:AL009126] subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis from 2000171to 2207900.] [LE:137942] [RE:138166]				
ORF Name	$\frac{\text{NTID}}{\text{NTID}} \frac{\text{AAID}}{\text{Length}} \frac{\text{NT}}{\text{Length}} \frac{\text{AA}}{\text{score}} \text{probability}$				
A17503000982_2381885_c1_988	316 4088 1272 423 1066 8.1e-108				
homolog ycsG] [GN:ycsG] [OR:Bacillu [LN:BSUB0003] [AC:Z99106:AL009126] [[DB:genpept-bct1] [DE:Bacillus subti	39895] [PN:branched chain amino acids transporter us subtilis] [DB:pir2] >gp:[GI:e1182373:g2632707] [GN:ycsG] [FN:unknown] [OR:Bacillus subtilis] ilis complete genome (section 3 of 21): from 402751 ycsH; similar to branched] [LE:54666] [RE:55826]				

ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	score	probabili	ty
A17503000982_23836052_f2_497	317	4089	849	282	392	2.1e-36	
Description					-		
sp:[LN:PROI_BACSU] [AC:P54552] [GN:Y [DE:PYRROLINE-5-CARBOXYLATE REDUCTAS pir:[LN:G69964] [AC:G69964] [PN:py [GN:yqj0] [CL:pyrroline-5-carboxyla pgp:[GI:d1013301:g1303966] [LN:BACJH subtilis] [SR:Bacillus subtilis (str [DE:Bacillus subtilis DNA, 283 Kb re [RE:246499] [DI:complement] pgp:[GI:[AC:Z99116:AL009126] [GN:yqj0] [FN:u [DE:Bacillus subtilis complete genom [NT:similar to pyrroline-5-carboxyla [DI:direct]	E HOMOLO rroline te reduc 642] [Ac ain:JH6 gion con e118564: nknown] e (sect:	OG 2] -5-carl ctase] C:D844 42(trpo ntainin 9:g2634 [OR:Ba ion 13	[SP:P54 coxylate [OR:Ba 32:D823 C2 Pheang skin 4815] [acillus of 21)	te reductions for the second s	DB:sw. ictase s subt: PN:Yqj(IA] [DI ent.] DB0013 .lis]	homolog yqjilis] [DB:pi D] [OR:Bacil B:genpept-bc [LE:245663] [DB:genpept-261to 261373	r2] lus t1] bct1]
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probabilit	ΞY
A17503000982_23850302_c3_1369	318	4090	2694	897	2695	1.9e-280	
Description sp:[LN:SYA_BACSU] [AC:O34526] [GN:AL							
[DE:ALANYL-TRNA SYNTHETASE, (ALANINE pir:[LN:A69584] [AC:A69584] [PN:al GN:alas] [CL:alaninetrnA ligase] pgp:[GI:e1183970:g2635186] [LN:BSUB0 synthetase] [GN:alas] [OR:Bacillus signer [GE:Bacillus subtilis complete genom [SP:O34526] [LE:197994] [RE:200630] [LN:BSUB0015] [AC:Z99118:AL009126] [subtilis] [DB:genpept-bct1] [EC:6.1.15 of 21): from 2795131to 3013540.]	TRNA 1 anine1 [OR:Bac 014] [Ac ubtilis] e (sect: [DI:comp PN:alany 1.7] [DI	LIGASE) tRNA li cillus C:Z9911 [DB:g ion 14 plement yl-tRNA E:Bacil	(ALAR igase, subtil 17:AL00 genpept of 21) c] >gp: A synth llus su	alaS:a alaS:a is] [E 99126] :-bct1] : from [GI:e1 actase]	P:034! C:6.1 [PN:a] [EC:6] 25994 183988 [GN:a]	526] [DB:swing the synthes and synthes and synthes and synthes and synthes are synthes and synthes are synthesis and synthesis are synthesis are synthesis.	tase] r2] 0.] cillus (section
ORF Name	NTID	AAID	<u>NT</u> Length	AA	score	probabilit	<u> </u>
A17503000982_23884692_c3_1415	319		<u> 1866 </u>	<u>neng cii</u>	217	7.5e-18	
Description Sp: [LN:YQHY_BACSU] [AC:P54519] [GN:YGKD PROTEIN IN ACCC-FOLD INTERGENIC R. [AC:E69960] [PN:conserved hypothetic [DB:pir2] >gp: [GI:d1013249:g1303914] [OR:Bacillus subtilis] [SR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis [LE:189273] [RE:189680] [DI:direct] [AC:Z99116:AL009126] [GN:yqhY] [FN:under subtilis complete genome subtilis complete genome subtilis complete genome subtilis complete genome subtilis subtilis complete genome subtilis complete genome subtilis subtilis complete genome subtilis subtilis complete genome subtilis su	QHY] [OI EGION] cal prot [LN:BAG subtilis lis DNA, >gp:[GI: nknown] e (secti	R:BACII [SP:P54 cein yo CJH642] s (stra 283 F :e11857 [OR:Ba	LLUS SU 1519] [G 1hY] [G 1n:JH6 Cb regi 701:g26 acillus of 21)	BTILIS DB:swi N:yqhY 84432: 42(trp on con 34867] subti : from] [DE:ssprot] [OF D82370 C2 Phetainir [LN:F] lis]	HYPOTHETICAL >pir:[LN:I :Bacillus su [PN:YqhY] (PN:YqhY] (PN:YqhY) (PN:YqhY)	E69960] ubtilis] ent.]
ORF Name	NTID	AAID	<u>NT</u> Length:	<u>AA</u> Length	score	probabilit	ΞY
AI7503000982 23912502 f2 605	320		1138				

Description NO-HIT

ORF Name A17503000982_23964011_t2_352	NTID	AAID NT AA score probability 4093 195 64				
Description						
NO-HIT						
ORF Name	NTID	AAID <u>NT AA</u> score probability				
AI7503000982_24042212_c3_1403	322	4094 498 165 72 0.034				
PROTEIN 3 PRECURSOR] [SP:P25955] [DB [AC:D30338:A35133:D69603] [PN:exoge 3] [GN:comGC] [OR:Bacillus subtilis [AC:M29691:M22854] [GN:comG3] [OR:Ba (clone: pED4) DNA] [DB:genpept-bct1] comG-(1,2,3,4,5,6,and 7) proteins in [DI:direct] >gp:[GI:d1013214:g130387 [OR:Bacillus subtilis] [SR:Bacillus [DB:genpept-bct1] [DE:Bacillus subtilis:] [SR:Bacillus subtilis:] [SR:Bacillus subtilis:] [AC:Z99116:AL009126] [GN:comGC] [FN:	:swissp: nous DNA] [DB:p: cillus a [DE:Bac comG ope 9] [LN:l subtilia lis DNA >gp:[GI exogenou	A-binding protein comGC:comG operon protein ir2] >gp:[GI:g142708] [LN:BACCOMGA] subtilis] [SR:Bacillus subtilis (strain 168) cillus subtilis (clone pED4) eron, complete cds.] [LE:3405] [RE:3701] BACJH642] [AC:D84432:D82370] [PN:ComGC] s (strain:JH642(trpC2 PheA1)) DNA] 283 Kb region containing skin element.] cel185739:g2634905] [LN:BSUB0013] us DNA-binding (competence)] [OR:Bacillus otilis complete genome (section 13 of 21):				
ORF Name AI7503000982 24071068 fl 186	NTID	AAID NT AA score probability LengthLength 308 1.7e-27				
Description						
sp:[LN:YQHL_BACSU] [AC:P54510] [GN:YQHL] [OR:BACILLUS SUBTILIS] [DE:HYPOTHETICAL 14.6 KD PROTEIN IN GCVT-SPOIIIAA INTERGENIC REGION] [SP:P54510] [DB:swissprot] >pir:[LN:C69959] [AC:C69959] [PN:glpE protein homolog yqhL] [GN:yqhL] [CL:glpE protein] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:d1013228:g1303893] [LN:BACJH642] [AC:D84432:D82370] [PN:YqhL] [OR:Bacillus subtilis] [SR:Bacillus subtilis (strain:JH642(trpC2 PheA1)) DNA] [DB:genpept-bct1] [DE:Bacillus subtilis DNA, 283 Kb region containing skin element.] [LE:174230] [RE:174610] [DI:complement] >gp:[GI:e1185722:g2634888] [LN:BSUB0013] [AC:Z99116:AL009126] [GN:yqhL] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 13 of 21): from 2395261to 2613730.] [NT:similar to hypothetical proteins] [SP:P54510] [LE:149011] [RE:149391] [DI:direct]						
ORF Name	NTID	AAID LengthLength score probability				
AI7503000982_24094090_c1_986	324	4096 957 318 927 4.4e-93				
<pre>subtilis] [DB:pir2] >gp:[GI:e1183965 [GN:yrrN] [FN:unknown] [OR:Bacillus et al., 1985]</pre>	:g263518 subtilis	homolog yrrN] [GN:yrrN] [OR:Bacillus B1] [LN:BSUB0014] [AC:Z99117:AL009126] B] [DB:genpept-bct1] [DE:Bacillus subtilis B9451to 2812870.] [NT:similar to protease]				

[LE:193967] [RE:194896] [DI:complement]

ORF Name LengthLength

AI7503000982_24100715_c3_1422

probability

1035

344

Description

sp:[LN:ODBA_BACSU] [AC:P37940] [GN:BFMBAA:BFMB1A] [OR:BACILLUS SUBTILIS] [EC:1.2.4.4] [DE:(BCKDH E1-ALPHA)] [SP:P37940] [DB:swissprot] >pir:[LN:C69593] [AC:C69593:S32486] [PN:3-methyl-2-oxobutanoate dehydrogenase (lipoamide), E1 alpha chain bfmBAA:branched-chain alpha-oxo acid dehydrogenase E1 alpha chain] [GN:bfmBAA] [CL:pyruvate dehydrogenase (lipoamide) alpha chain: thiamine pyrophosphate-binding domain homology] [OR:Bacillus subtilis] [EC: 1.2.4.4] [DB:pir2] >qp:[GI:q142611] [LN:BACBRANCH] [AC:M97391:M96937] [PN:branched chain alpha-keto acid dehydrogenase] [OR:Bacillus subtilis] [SR:Bacillus subtilis DNA] [DB:genpept-bct1] [DE:Bacillus subtilis branched chain alpha-keto acid dehydrogenaseE1-alpha, branched chain alpha-keto acid dehydrogenase El-beta, andbranched chain alpha-keto acid dehydrogenase E2, complete cds.] [LE:216] [RE:1208] [DI:direct] >gp:[GI:d1013277:g1303942] [LN:BACJH642] [AC:D84432:D82370] [PN:BfmBAA] [OR:Bacillus subtilis] [SR:Bacillus subtilis (strain:JH642(trpC2 PheA1)) DNA] [DB:genpept-bct1] [DE:Bacillus subtilis DNA, 283 Kb region containing skin element.] [LE:219562] [RE:220554] [DI:direct] >gp:[GI:e1185673:g2634839] [LN:BSUB0013] [AC:Z99116:AL009126] [PN:branched-chain alpha-keto acid dehydrogenase E1] [GN:bfmBAA] [OR:Bacillus subtilis] [DB:genpept-bct1] [EC:1.2.4.4] [DE:Bacillus subtilis complete genome (section 13 of 21): from 2395261to 2613730.] [NT:alternate gene name: bfmB1a] [SP:P37940] [LE:103065] [RE:104057] [DI:complement]

ORF Name	NTID	AAID	NT AA LengthLength score probability
AI7503000982_24104702_c1_1109	326	4098	3061 10,20 618 1.1e-54
Description			· .
gp:[GI:g1041785] [LN:PYU36927] [AC:U invasion and possible binding] [OR:P			optry protein] [FN:erythrocyte lii] [DB:genpept-inv1] [DE:Plasmodium
yoelii rhoptry protein gene, partial	cds.]	[LE:<1] [RE:7206] [DI:direct]
ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000982_24117777_f3_858	327	4099	174 57
Description			
NO-HIT			
ORF Name	NTID	AAID	NT <u>AA</u> LengthLength
A17503000982_24225000_c1_1152	328	4100	651 216 365 1.6e-33

Description

gp:[GI:g2194195] [LN:SGU61158] [AC:U61158] [PN:GdmF] [GN:gdmF] [OR:Staphylococcus gallinarum [DB:genpept-bct1] [DE:Staphylococcus gallinarum Tue3928 GdmF (qdmF), putative membraneprotein (gdmH), ABC transporter (gdmT), and antibiotic galliderminprecursor (gdmA) genes, complete cds, putative membrane protein(gdmE) and modifying enzyme (gdmB) genes, partial cds.] [NT:proposed ABC transporter subunit (ATP-binding] [LE:179] [RE:874] [DI:complement]

ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	score	probability
A17503000982_24226412_c3_1436	329	4101	570	189		3.7e-41
Description			J [J L	
sp:[LN:YQKG_BACSU] [AC:P54570] [GN:YGKD PROTEIN IN GLNQ-ANSR INTERGENIC RIAC:A69967] [PN:conserved hypothetic domain homology] [OR:Bacillus subtiling [LN:BACJH642] [AC:D84432:D82370] [PN (strain:JH642(trpC2 PheA1)) DNA] [DB region containing skin element.] [LE >gp:[GI:e1185630:g2634796] [LN:BSUB06] [OR:Bacillus subtilis] [DB:genpept-b6] of 21): from 2395261to 2613730.] [LE:62478] [RE:63035] [DI:complement]	EGION] cal pro is] [DB :YqkG] :genpep :260584 D13] [A ct1] [D	[SP:P56 tein yo :pir2] [OR:Bao t-bct1] [RE:2 C:Z991:	4570] qkG] [0 >gp:[0 cillus] [DE:F 261141] 16:AL00 llus su	[DB:swi GN:yqkG GI:d101 subtill Gacillu [DI:d (9126] ubtilis	issprot G] [CI .3319:g .is] [S is subt direct] [GN:yq s compl	pir:[LN:A69967] syffH protein:mutT [1303984] R:Bacillus subtilis cilis DNA, 283 Kb [kG] [FN:unknown] ete genome (section
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000982_24226635_£2_499	330	4102	156	51]	
Description No. W.T.						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000982_24228452_f3_942	331	4103][138	45	:]	
Description		L	J []		J	
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
AI7503000982_24229515_c3_1423	332	4104	1170	389		1.1e-98
Description sp:[LN:YQJE_BACSU] [AC:P54542] [GN:YQKD PROTEIN IN GLNQ-ANSR INTERGENIC RI [AC:E69963] [PN:tripeptidase homolog >gp:[GI:d1013291:g1303956] [LN:BACJHG subtilis] [SR:Bacillus subtilis (stra [DE:Bacillus subtilis DNA, 283 Kb reg [RE:233971] [DI:direct] >gp:[GI:e1185] [GN:yqjE] [FN:unknown] [OR:Bacillus scomplete genome (section 13 of 21): ftripeptidase] [SP:P54542] [LE:89648]	EGION] y yqjE] 642] [A ain:JH6 gion con 6659:g20 subtilis from 233	[SP:P54 [GN:y0 C:D8443 42(trp0 ntainir 534825] s] [DB:	1542] [1jE] [132:D823 122 PheAng skin 1 [LN:E 1:genpep 1:26137	DB:swi OR:Bac 70] [P 1)) DN eleme SUB001 t-bct1	ssprot illus N:YqjE [A] [DB nt.] [AC] [DE: NT:sim	<pre>] >pir:[LN:E69963] subtilis] [DB:pir2]] [OR:Bacillus :genpept-bct1] LE:232856] :Z99116:AL009126] Bacillus subtilis</pre>
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000982_24229805_c3_1376	333	4105	1386	461	1133	6.4e-115
Description				/1- d - 1 d		
pir:[LN:A69581] [AC:A69581] [PN:acet accC] [GN:accC] [CL:biotin carboxyla subtilis] [DB:pir2] >gp:[GI:d1013248:[PN:YqhX] [OR:Bacillus subtilis] [SR:[DB:genpept-bct1] [DE:Bacillus subtilic] [LE:187900] [RE:189252] [DI:direct] > [AC:Z99116:AL009126] [PN:acetyl-CoA of [FN:long-chain fatty acid biosynthesis [EC:6.4.1.2] [DE:Bacillus subtilis county [Coanglement]	ase:biot g130393 Bacillu is DNA gp:[GI carboxyl s] [OR	in car L3] [LN us subt 283 K el1857 Lase su Bacill genome	Tboxyla I:BACJH Tilis (Tb regi T02:g26 Tbunit Tus sub	se hom [642] [strain on con 34868] (bioti tilis] ion 13	ology] AC:D84 :JH642 tainin [LN:B n] [GN [DB:g of 21	[OR:Bacillus 432:D82370] (trpC2 PheA1)) DNA] g skin element.] SUB0013] :accC] enpept-bct1]): from 2395261to

ORF Name A17503000982_24257658_f1_309 Description NO-HIT	NTID	<u>AAID</u> 4106	NT AA LengthLength 162 53	probability
ORF Name A17503000982_24260061_c3_1353 Description	<u>NTID</u>	<u>AAID</u> 4107	NT AA LengthLength score	probability
sp:[LN:YSXB_BACSU] [AC:P26942:Q45629] 12.3 KD PROTEIN IN RPLU-RPMA INTERGER [DB:swissprot] >pir:[LN:S18440] [AC:Sprotein ysxB] [GN:ysxB] [OR:Bacillus [LN:BSSPOIVFO] [AC:X59528:S61796] [OR [DE:B.subtilis spoIVFA, spoIVFB, L20] [LE:2270] [RE:2608] [DI:direct] >gp: [AC:Z99118:AL009126] [GN:ysxB] [FN:un [DE:Bacillus subtilis complete genome [NT:similar to hypothetical proteins]	NIC REGIS18440:Is subtil R:Bacill , orfX a [GI:e118 nknown] e (secti	ON (O) 021895 Lis] [1 Lus sub and L24 34044:0 [OR:Ba	RF X)] [SP:P26942:Q :B69987] [PN:conse DB:pir2] >gp:[GI:g4 otilis] [DB:genpept 4 genes.] [NT:ORF X g2635260] [LN:BSUB0 acillus subtilis] [of 21): from 27951	45629] rved hypothetical 0174] -bct1]] [SP:P26942] 015] DB:genpept-bct1] 31to 3013540.]
ORF Name A17503000982_24261692_c3_1518	<u>NTID</u>	<u>AAID</u> 4108	NT AA LengthLength score	probability 2.7e-100
Description pir: [LN:D69907] [AC:D69907] [PN:hyposubtilis] [DB:pir2] >gp:[GI:e1185410] [GN:yojO] [FN:unknown] [OR:Bacillus stomplete genome (section 11 of 21): ff [DI:complement] >gp:[GI:g3169331] [LN [FN:unknown] [OR:Bacillus subtilis] (yojA), YojB (yojB), YojC (yojC), YojYojH (yojH), YojI(yojI), YojJ (yojJ), YojN(yojH), and YojO (yojO) genes, colle:12366] [RE:14351] [DI:direct]	:g263433 subtilis Erom 200 N:AF0261 [DB:genr jD(yojD) , YojK (31] [Li 3] [DB: 00171tc .47] [i pept-bo , Yoji (YojK),	N:BSUB0011] [AC:Z99 genpept-bct1] [DE: 2207900.] [LE:110 AC:AF026147] [PN:Yo ct2] [DE:Bacillus s E (yojE), YojF (yoj YojL (yojL), YojM	114:AL009126] Bacillus subtilis 914] [RE:112899] jO] [GN:yojO] ubtilis YojA F), YojG (yojG), (yojM),
ORF Name A17503000982_24266502_t3_675 Description NO-HIT	NTID 337	<u>AAID</u> 4109	NT AA LengthLength	probability
ORF Name A17503000982_24274192_f2_386 Description	NTID 338	AAID 4110	NT AA LengthLength score	probability
sp:[LN:EBSB_ENTFA] [AC:P36921] [OR:EN [DE:CELL WALL ENZYME EBSB] [SP:P36921] [PN:ebsB protein] [CL:Enterococcus fa [DB:pir2] >gp:[GI:g388108] [LN:ENEEBS [OR:Enterococcus faecalis] [SR:Enterococcus faecalis] [DB:genpept-bct2] [DE:Enterococcus faeculatory, and dehydroquinase homologcomplete cds with repeat region.] [NT	l] [DB:s mecalis BA] [AC: pcoccus mecalis gue prot	wissprebsB p L23802 faecal pore f	rot] >pir:[LN:B4993] protein] [OR:Enteroon] [PN:cell wall end is (strain OG1SSp) forming, cell wall of [ebsA,ebsB,ebsC,and	P] [AC:B49939] COCCUS faecalis] Zyme] [GN:ebsB] DNA] enzyme, ebsD)genes,

ORF Name AI7503000982_24275927_f2_622 Description NO-HIT	NTID	<u>AAID</u>	NT AA LengthLength score	probability
ORF Name A17503000982_24306263_c1_1034 Description NO-HIT	NTID 340	AAID 4112	NT AA LengthLength 297 98	probability
ORF Name A17503000982_24410300_c3_1439 Description sp:[LN:RLUB_BACSU] [AC:P35159] [GN:RLUB_BCSU] [AC:P35159] [GN:RLUB_DE:(PSEUDOURIDYLATE SYNTHASE) (URACE) >pir:[LN:S45555] [AC:S45555:A69943]] [CL:conserved hypothetical protein >gp:[GI:g410137] [LN:BACDIA] [AC:L0991] (strain 168, sub_species Marburg) DN: serA region.] [NT:ORFX13] [LE:17440] [LN:BSUB0013] [AC:Z99116:AL009126] [GUB:genpept-bct1] [DE:Bacillus subtile 2395261to 2613730.] [NT:similar to hypothement]	IL HYDRO [PN:con HI1243] 228] [OH A] [DB:com [RE:183] GN:ypuL]	OLYASE nserve [OR: R:Baci genpep 129] [[FN: plete	LLUS SUBTILIS] [EC:)] [SP:P35159] [DB: d hypothetical prot Bacillus subtilis] llus subtilis] [SR: t-bct1] [DE:Bacillu DI:direct] >gp:[GI: unknown] [OR:Bacill genome (section 13	swissprot] ein ypuL] [GN:ypuL [DB:pir2] Bacillus subtilis s subtilis spoVA to e1185585:g2634751] us subtilis] of 21): from
ORF Name A17503000982_24470317_c1_1011 Description sp:[LN:YQFA_BACSU] [AC:P54466] [GN:YQKD PROTEIN IN RPSU-PHOH INTEREGENIC IN [AC:A69953] [PN:hypothetical protein >gp:[GI:d1013152:g1303817] [LN:BACJHG subtilis] [SR:Bacillus subtilis (strate) [DE:Bacillus subtilis DNA, 283 Kb reg [RE:102156] [DI:direct] >gp:[GI:e1185] [GN:yqfA] [FN:unknown] [OR:Bacillus subtilis complete genome (section 14 of 21): [RE:18269] [DI:complement]	REGION] n yqfA] 642] [AG ain:JH64 gion cor 3768:g26 subtilis	[SP:P [GN:y C:D844 12(trp ntaini: 534984 5] [DB	LLUS SUBTILIS] [DE: 54466] [DB:swisspro qfA] [OR:Bacillus 32:D82370] [PN:YqfA C2 PheA1)) DNA] [DB ng skin element.] [] [LN:BSUB0014] [AC:genpept-bct1] [DE:	t] >pir:[LN:A69953] subtilis] [DB:pir2]] [OR:Bacillus :genpept-bct1] LE:101161] :Z99117:AL009126] Bacillus subtilis
ORF Name A17503000982_24480275_c3_1499 Description pir:[LN:A69653] [AC:A69653] [PN:tran [OR:Bacillus subtilis] [DB:pir2] >gp [AC:Z99107:AL009126] [PN:transmembran [DB:genpept-bct1] [DE:Bacillus subtil to813890.] [LE:179900] [RE:180856] [I	:[GI:e11 ne lipop lis comp	182690 protein plete (poprotein lplB] [GN:g2633024] [LN:BSUB n] [GN:lplB] [OR:Ba	0004] cillus subtilis]

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000982_24486330_c1_980	344	4116	693	230	321	7.2e-29
Description pir: [LN:H69978] [AC:H69978] [PN:con [CL:tetratricopeptide repeat homolog >gp: [GI:e1183978:g2635194] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-b 14 of 21): from 2599451to 2812870.] [RE:209147] [DI:complement] >gp:[GI: [AC:Z99118:AL009126] [GN:yrrB] [FN:u [DE:Bacillus subtilis complete genom [NT:similar to hypothetical proteins	y] [OR:1 014] [A ct1] [D [NT:sim: e1183990 nknown] e (sect:	Bacilly C:Z991 E:Bacil ilar to 6:g263! [OR:Ba ion 15	us subt 17:ALOC llus su o hypot 5212] acillus of 21)	cilis] 09126] ubtilis chetica [LN:BSU s subti : from	[DB:pi [GN:yr comp] l prot B0015] lis]	rr2] rrB] [FN:unknown] ete genome (section eins] [LE:208527] DB:genpept-bct1] 31to 3013540.]
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
AI7503000982_24489062_f1_1	345	4117	126	41	72	0.017
Description pir: [LN:D70083] [AC:D70083] [PN:hyp subtilis] [DB:pir2] >gp: [GI:e1184645 [GN:yxzC] [FN:unknown] [OR:Bacillus complete genome (section 21 of 21): [DI:complement]	:g263646 subtilis	66] [L1 s] [DB:	N:BSUBC genpep	0021] [. ot-bct1	AC:Z99	124:AL009126] Bacillus subtilis
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000982_24511676_c1_1140	346	4118	246	81	251	4.3e-21
Description sp:[LN:ODO2_BACSU] [AC:P16263] [GN:ODE:DEHYDROGENASE COMPLEX, (E2)] [SPENCE:B32879:F69668] [PN:dihydrolipoate dehydrogenase complex E2 component of [GN:odhB] [CL:dihydrolipoamide acety [OR:Bacillus subtilis] [EC:2.3.1.61] [AC:M27141] [OR:Bacillus subtilis] [ED:B.subtilis 2-oxoglutarate dehydrotranssuccinylase (odhB) gene, complete (odhB; EC] [LE:899] [RE:2152] [DI:di:[AC:Z99114:AL009126] [PN:2-oxoglutarasubtilis] [DB:genpept-bct1] [EC:2.3.1] (section 11 of 21): from 2000171to 2:[LE:106590] [RE:107843] [DI:complement]	:P16263] mide S-s dhB: dil yltransf [DB:pin SR:B.suk ogenase te cds.] rect] >9 ate deh 1.61] [I 207900.]] [DB:ssucciny hydroliferase: r2] >gg btilis (odhA)] [NT:cogp:[GI: ydrogen DE:Baci	swisspr vltrans ipoamid : lipoy D:[GI:g (strai gene dihydro :e11854 nase co	ot] >p ferase le trans 1/biot 143268 n 3G18 3' end lipoam 08:g26 mplex] ubtilis	ir:[LN; , odhEssucci in-bin] [LN:) DNA] , andd ide tr 34329] [GN:os comp	E:B32879] E:2-oxoglutarate nylase odhB] ding homology] BACODHAB] [DB:genpept-bct1] ihydrolipoamide anssuccinylase [LN:BSUB0011] dhB] [OR:Bacillus lete genome
ORF Name	NTID		<u>NT</u> Length	rend cu	score	probability
AI7503000982_24610877_c1_1035 Description	347	4119	519	172	226	8.4e-19
sp:[LN:AROK_LACLA] [AC:P43906] [GN:AI [SR:,SUBSPLACTIS:STREPTOCOCCUS LACTIS [SP:P43906] [DB:swissprot] >pir:[LN:S [CL:shikimate kinase:shikimate kinase [DB:pir2] >gp:[GI:g683584] [LN:LLTYRA [OR:Lactococcus lactis] [DB:genpept-B	S] [EC:2 S52581] e homolo APH] [AC	2.7.1.7 [AC:S5 ogy] [C C:X7841	71] [DE 52581] DR:Lact .3] [PN	SHIKIN: [PN:sl ococcus shikin:	MATE K nikima s lact nate k	te kinase,] is] [EC:2.7.1.71] inase] [GN:aroK]

and pheA genes.] [SP:P43906] [LE:2605] [RE:3093] [DI:direct]

ORF Name	NTID	AAID LengthLength score probability
A17503000982_24615915_c3_1522	348	4120 240 79 275 5.4e-24
	lococcu] [PN:CspC] [GN:cspC] [OR:Staphylococcus s aureus CspC (cspC) gene, complete cds.] E:444] [RE:644] [DI:direct]
ORF Name	NTID	AAID NT AA score probability
A17503000982_24617177_c3_1433	349	4121 129 42
Description NO-HIT		
ORF Name	NTID	AAID NT AA score probability
AI7503000982_24640937_c2_1229	350	4122 498 165
Description NO-HIT		
ORF Name	NTID	AAID NT AA score probability
AI7503000982_24647187_c1_1038	351	4123 126 41 98 8.8e-05
haemolyticus] [SR:Staphylococcus hae [DE:Staphylococcus haemolyticus IS12 [LE:394] [RE:1083] [DI:complement] > [OR:Staphylococcus aureus] [SR:Staph [DB:genpept-bct1] [DE:Staphylococcus	molytic 72 ORF1 gp:[GI: ylococc aureus	and ORF2 genes, completecds.] [NT:ORF2] g295162] [LN:STAMECRA] [AC:L14017]
ORF Name	NTID	AAID NT AA score probability
AI7503000982_24650016_c3_1478	352	4124 417 138 155 2.8e-11
KD PROTEIN IN COTD-KDUD INTERGENIC R [AC:E69941] [PN:hypothetical protei >gp:[GI:g1146182] [LN:BACPONAYPP] [A subtilis] [DB:genpept-bct1] [DE:Baci genes, ypqAE genes, yprAB genes, cotD gene, kduDI genes, kdgRKAT genes, yp [RE:12488] [DI:direct] >gp:[GI:e1183 [GN:ypsB] [FN:unknown] [OR:Bacillus	EGION] n ypsB] C:L47833 llus sul gene, wA gene 664:g263 subtilis	R:BACILLUS SUBTILIS] [DE:HYPOTHETICAL 11.6 [SP:P50839] [DB:swissprot] >pir:[LN:E69941] [GN:ypsB] [OR:Bacillus subtilis] [DB:pir2] 8] [GN:ypsB] [FN:hypothetical] [OR:Bacillus btilis (clone YAC15-6B) ponA gene, yppBCDEFG ypsABC genes, rnaP gene, yptAgene, ypuA, completecds's.] [NT:putative] [LE:12192] 34637] [LN:BSUB0012] [AC:Z99115:AL009126] s] [DB:genpept-bct1] [DE:Bacillus subtilis 95541to 2409220.] [SP:P50839] [LE:135481]
ORF Name	NTID	AAID NT AA score probability
A17503000982_24650252_f3_731 Description gp:[GI:e1357086:g3955030] [LN:SAU177	353 95] [AC	4125 633 210 970 1.2e-97
	pt-bct1]	[DE:Staphylococcus aureus prfA, pbp2

<u>ORF Name</u>

NTID AAID NT AA LengthLength score probability

A17503000982_24658562_f3_926

354 | 4126 | 1287 | 428 | 1475 | 3.7e-151

pir:[LN:D69981] [AC:D69981] [PN:conserved hypothetical protein yrvN] [GN:yrvN] [CL:Haemophilus influenzae conserved hypothetical protein HI1590] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1183983:g2635199] [LN:BSUB0014] [AC:Z99117:AL009126] [GN:yrvN] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 14 of 21): from 2599451to 2812870.] [NT:similar to hypothetical proteins] [LE:212152] [RE:213417] [DI:direct] >gp:[GI:e1184001:g2635217] [LN:BSUB0015] [AC:Z99118:AL009126] [GN:yrvN] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 15 of 21): from 2795131to 3013540.] [NT:similar to hypothetical proteins] [LE:16472] [RE:17737] [DI:direct]

 ORF Name
 NTID
 AAID
 NT AA Length Length
 Score
 probability

 A17503000982_24797827_c3_1514
 355
 4127
 1857
 618
 1242
 1.8e-126

Description

sp:[LN:ODO1_BACSU] [AC:P23129] [GN:ODHA:CITK] [OR:BACILLUS SUBTILIS] [EC:1.2.4.2]
[DE:KETOGLUTARATE DEHYDROGENASE)] [SP:P23129] [DB:swissprot] >pir:[LN:A32879]
[AC:S25295:A32879:E69668:S14544] [PN:oxoglutarate dehydrogenase (lipoamide),] [GN:odhA]
] [CL:oxoglutarate dehydrogenase (lipoamide):thiamine pyrophosphate-binding domain
homology] [OR:Bacillus subtilis] [EC:1.2.4.2] [DB:pir2] >gp:[GI:g40003] [LN:BSODHA]
[AC:X54805:S43328] [PN:oxoglutarate dehydrogenase (NADP+)] [GN:odhA] [OR:Bacillus
subtilis] [DB:genpept-bct1] [DE:B. subtilis odhA gene for 2-oxoglutarate
dehydrogenase.] [SP:P23129] [LE:291] [RE:3104] [DI:direct] >gp:[GI:e1185409:g2634330]
[LN:BSUB0011] [AC:Z99114:AL009126] [PN:2-oxoglutarate dehydrogenase (E1 subunit)]
[GN:odhA] [OR:Bacillus subtilis] [DB:genpept-bct1] [EC:1.2.4.2] [DE:Bacillus subtilis
complete genome (section 11 of 21): from 2000171to 2207900.] [NT:alternate gene name:
citK] [SP:P23129] [LE:107873] [RE:110686] [DI:complement]

 ORF Name
 NTID
 AAID
 NT AA Length Length
 score
 probability

 A17503000982_24800461_c1_1138
 356
 4128
 231
 76
 235
 1.7e-18

Description

sp:[LN:ODO1_BACSU] [AC:P23129] [GN:ODHA:CITK] [OR:BACILLUS SUBTILIS] [EC:1.2.4.2]
[DE:KETOGLUTARATE DEHYDROGENASE)] [SP:P23129] [DB:swissprot] >pir:[LN:A32879]
[AC:S25295:A32879:E69668:S14544] [PN:oxoglutarate dehydrogenase (lipoamide),] [GN:odhA]
[CL:oxoglutarate dehydrogenase (lipoamide):thiamine pyrophosphate-binding domain homology] [OR:Bacillus subtilis] [EC:1.2.4.2] [DB:pir2] >gp:[GI:g40003] [LN:BSODHA]
[AC:X54805:S43328] [PN:oxoglutarate dehydrogenase (NADP+)] [GN:odhA] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:B. subtilis odhA gene for 2-oxoglutarate dehydrogenase.] [SP:P23129] [LE:291] [RE:3104] [DI:direct] >gp:[GI:e1185409:g2634330] [LN:BSUB0011] [AC:Z99114:AL009126] [PN:2-oxoglutarate dehydrogenase (E1 subunit)]
[GN:odhA] [OR:Bacillus subtilis] [DB:genpept-bct1] [EC:1.2.4.2] [DE:Bacillus subtilis complete genome (section 11 of 21): from 2000171to 2207900.] [NT:alternate gene name: citK] [SP:P23129] [LE:107873] [RE:110686] [DI:complement]

ORF Name	MIID	Length Length Scote Probability
A17503000982_24807790_c1_1048	357	4129 909 302 687 1.2e-67
Description sp:[LN:ISPA_BACST] [AC:Q08291] [OR:B SYNTHASE)] [SP:Q08291] [DB:swissprot [PN:geranyltranstransferase,:farnesy [CL:geranyltranstransferase] [OR:Bac >gp:[GI:d1003054:g391610] [LN:BACFDP	ACILLUS] >pir: -1-diphos -illus si -S] [AC:1 :Bacillus	STEAROTHERMOPHILUS] [EC:2.5.1.10] [DE:(FPP [LN:JX0257] [AC:JX0257] sphate synthase] tearothermophilus] [EC:2.5.1.10] [DB:pir2] D13293] [PN:farnesyl diphosphate synthase] as stearothermophilus (strain:ATCC10149) stearothermophilus DNA for farnesyl
		ex-determining region hypothetical protein c] [DB:pir2]
ORF Name AI7503000982_24886562_f3_771 Description NO-HIT	<u>NTID</u>	AAID LengthLength score probability 4131 123 40
[OR:Bacillus subtilis] [DB:pir2] >gp [AC:Z99117:AL009126] [GN:yrrC] [FN:u [DE:Bacillus subtilis complete genom [NT:similar to conjugation transfer >gp:[GI:e1183995:g2635211] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-b	:[GI:e1: nknown] e (sect: protein] 015] [AG ct1] [DI [NT:sim	AAID NT AA LengthLength AAID LengthLength AAID LengthLength AIII LengthLength AIIII LengthLength AIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIII
A17503000982_2542188_c1_1026 Description NO-HIT	361	4133 270 89
ORF Name A17503000982_25423425_f2_337 Description NO-HIT	NTID 362	AAID NT AA score probability LengthLength 69
ORF Name A17503000982_25428378_t2_336 Description NO-HIT	NTID 363	AAID NT AA score probability LengthLength 75

Sescription Sep: [LN:RUVA_BACSU] [AC:005392] [GN:RUVA] [OR:BACILLUS SUBTILIS] [DE:PROBABLE HOLLIDAY DUNCTION DNA HELICASE RUVA] [SP:005392] [DB:swissprot] >pir: [LN:E69702] [AC:E69702] [PN:Holliday junction DNA helicase ruvA] [GN:ruvA] [CL:holliday junction DNA helicase ruvA] [GN:ruvA] [GN:Bacillus subtilis] [DB:pir2] >gp: [GI:e1184023:g2635239] [LN:BSUB0015] [AC:Z99118:AL009126] [PN:Holliday junction DNA helicase] [GN:ruvA] [OR:Bacillus subtilis] [DB:genpept-bctl] [DE:Bacillus subtilis complete genome (section 15 of 21): from 2795131to 3013540.] [SP:005392] [LE:40296] [RE:40901] [DI:complement] Some
Sp:[LN:RUVA_BACSU] [AC:005392] [GN:RUVA] [OR:BACILLUS SUBTILIS] [DE:PROBABLE HOLLIDAY JUNCTION DNA HELICASE RUVA] [SP:005392] [DB:swissprot] >pir:[LN:E69702] [AC:E69702] [PN:Holliday junction DNA helicase ruvA] [GN:ruvA] [CL:holliday junction DNA helicase ruvA] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1184023:g2635239] [LN:BSUB0015] [AC:Z99118:AL009126] [PN:Holliday junction DNA helicase] [GN:ruvA] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 15 of 21): from 2795131to 3013540.] [SP:005392] [LE:40296] [RE:40901] [DI:complement] DRF Name NTID AAID LengthLength Description pir:[LN:G69728] [AC:G69728] [PN:uridine kinase udk] [GN:udk] [CL:uridine kinase] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1183963:g2635179] [LN:BSUB0014] [AC:Z99117:AL009126] [PN:uridine kinase] [GN:udk] [FN:pyrimidine salvage] [OR:Bacillus subtilis] [DB:genpept-bct1] [EC:2.7.1.48] [DE:Bacillus subtilis complete genome (section 14 of 21): from 2599451to 2812870.] [LE:192038] [RE:192673] [DI:complement]
JUNCTION DNA HELICASE RUVA] [SP:005392] [DB:swissprot] >pir:[LN:E69702] [AC:E69702] [PN:Holliday junction DNA helicase ruvA] [GN:ruvA] [CL:holliday junction DNA helicase ruvA] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1184023:g2635239] [LN:BSUB0015] [AC:Z99118:AL009126] [PN:Holliday junction DNA helicase] [GN:ruvA] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 15 of 21): from 2795131to 3013540.] [SP:005392] [LE:40296] [RE:40901] [DI:complement] DRF Name
DESCRIPTION AID LengthLength LengthLength AITS Score probability AITS Description pir: [LN:G69728] [AC:G69728] [PN:uridine kinase udk] [GN:udk] [CL:uridine kinase] [OR:Bacillus subtilis] [DB:pir2] >gp: [GI:e1183963:g2635179] [LN:BSUB0014] [AC:Z99117:AL009126] [PN:uridine kinase] [GN:udk] [FN:pyrimidine salvage] [OR:Bacillus subtilis] [DB:genpept-bct1] [EC:2.7.1.48] [DE:Bacillus subtilis complete genome (section 14 of 21): from 2599451to 2812870.] [LE:192038] [RE:192673] [DI:complement] ORF Name NTID AAID LengthLength Core probability AAID LengthLength LengthLength AAID LengthLength LengthLength LengthLength
Description pir: [LN:G69728] [AC:G69728] [PN:uridine kinase udk] [GN:udk] [CL:uridine kinase] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1183963:g2635179] [LN:BSUB0014] [AC:Z99117:AL009126] [PN:uridine kinase] [GN:udk] [FN:pyrimidine salvage] [OR:Bacillus subtilis] [DB:genpept-bct1] [EC:2.7.1.48] [DE:Bacillus subtilis complete genome (section 14 of 21): from 2599451to 2812870.] [LE:192038] [RE:192673] [DI:complement] ORF Name NTID AAID NT AA LengthLength
pir: [LN:G69728] [AC:G69728] [PN:uridine kinase udk] [GN:udk] [CL:uridine kinase] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1183963:g2635179] [LN:BSUB0014] [AC:Z99117:AL009126] [PN:uridine kinase] [GN:udk] [FN:pyrimidine salvage] [OR:Bacillus subtilis] [DB:genpept-bct1] [EC:2.7.1.48] [DE:Bacillus subtilis complete genome (section 14 of 21): from 2599451to 2812870.] [LE:192038] [RE:192673] [DI:complement] ORF Name NTID AAID NT AA LengthLength
NTID AAID score probability LengthLength
A17503000982_25598818_c3_1420
Sp:[LN:RECN_BACSU] [AC:P17894:P19671] [GN:RECN] [OR:BACILLUS SUBTILIS] [DE:DNA REPAIR PROTEIN RECN (RECOMBINATION PROTEIN N)] [SP:P17894:P19671] [DB:swissprot] [PROTEIN RECN (RECOMBINATION PROTEIN N)] [SP:P17894:P19671] [DB:swissprot] [PR:CLN:B35128] [AC:B35128:PS0054:H69690] [PN:DNA repair and genetic recombination protein recN] [GN:recN] [CL:recN protein] [OR:Bacillus subtilis] [DB:pir2] [PN:RecN] [OR:Bacillus subtilis] [SR:Bacillus subtilis (strain:JH642(trpC2 PheA1)) DNA] [DB:genpept-bct1] [DE:Bacillus subtilis DNA, 283 Kb region containing skin element.] [LE:197319] [RE:199049] [DI:direct] >gp:[GI:g143402] [LN:BACRECN] [AC:M30297] [OR:Bacillus subtilis] [SR:B.subtilis (strain 168) DNA] [DB:genpept-bct1] [DE:B.subtilis recombination and sporulation protein (recN, spoIVB)genes, complete cds, arginine hydroximate resistance (ahrC) gene,3' end.] [NT:recombination protein (ttg start codon)] [LE:131] [RE:1861] [DI:direct] >gp:[GI:e1185692:g2634858] [LN:BSUB0013] [AC:Z99116:AL009126] [GN:recN] [FN:DNA repair and genetic recombination] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 13 of 21): from 2395261to 2613730.] [SP:P17894] [LE:124572] [RE:126302] [DI:complement]
NTID AAID NT AA Score probability A17503000982 25652217 c3 1470 367 4139 348 115

Description NO-HIT

ORF Name	NTID AAID NT Length	AA Length score probability
A17503000982_25672337_c1_1016	368 4140 780	259 371 3.6e-34
Description		
sp:[LN:YQXN_BACSU] [AC:P42095] [GN:Y [SP:P42095] [DB:swissprot] >pir:[LN:protein yqxN] [GN:yqxN] [OR:Bacillu [LN:BACJH642] [AC:D84432:D82370] [PN (strain:JH642(trpC2 PheA1)) DNA] [DB region containing skin element.] [LE >gp:[GI:e1185794:g2634960] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-b13 of 21): from 2395261to 2613730.] hypothetical] [SP:P42095] [LE:212962 >gp:[GI:e1183758:g2634974] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-b14 of 21): from 2599451to 2812870.] hypothetical] [SP:P42095] [LE:8772]	[AC:H69968] [AC:H69968] subtilis] [DB:pir2 YqfI] [OR:Bacillus genpept-bct1] [DE:B 109891] [RE:110658] [AC:Z99116:AL00 t1] [DE:Bacillus su [RE:213729] [DI:co 14] [AC:Z99117:AL00 t1] [DE:Bacillus su [NT:alternate gene n [RE:213729] [DI:co 14] [AC:Z99117:AL00 t1] [DE:Bacillus su [NT:alternate gene n	[PN:conserved hypothetical] >gp:[GI:d1013162:g1303827] subtilis] [SR:Bacillus subtilis acillus subtilis DNA, 283 Kb [DI:direct] 9126] [GN:yqxN] [FN:unknown] btilis complete genome (section ame: yqfI; similar to mplement] 9126] [GN:yqxN] [FN:unknown] btilis complete genome (section ame: yqfI; similar to
ORF Name	NTID AAID NT Length	AA Length score probability
A17503000982_25680218_c2_1201	369 4141 726	241 425 6.8e-40
Description		
KD PROTEIN IN RPSU-PHOH INTEREGENIC [AC:H69952] [PN:hypothetical protei >gp:[GI:d1013151:g1303816] [LN:BACJH subtilis] [SR:Bacillus subtilis (str [DE:Bacillus subtilis DNA, 283 Kb re [RE:101139] [DI:direct] >gp:[GI:e118 [GN:yqeZ] [FN:unknown] [OR:Bacillus complete genome (section 14 of 21): [RE:19604] [DI:complement]	yqeZ] [GN:yqeZ] [GN:yqeZ] [42] [AC:D84432:D823 in:JH642(trpC2 PheA ion containing skin 769:g2634985] [LN:B ubtilis] [DB:genpep	OR:Bacillus subtilis] [DB:pir2] 70] [PN:YqeZ] [OR:Bacillus 1)) DNA] [DB:genpept-bct1] element.] [LE:99826] SUB0014] [AC:Z99117:AL009126] t-bct1] [DE:Bacillus subtilis
ORF Name	NTID AAID NT Lengthl	AA Length score probability
AI7503000982_26058467_c1_1064	370 4142 213	70 119 9.1e-07
Description gp:[GI:g861340] [LN:CELE04F6] [AC:U2 [SR:Caenorhabditis elegans strain=Br elegans cosmid E04F6.] [NT:similar t [RE:20659:20814:21152] [DI:complemen	stol N2] [DB:genpep ribitol dehydrogen	t-inv1] [DE:Caenorhabditis
ORF Name	NTID AAID NT LengthI	AA Length score probability
AI7503000982_26203942_c3_1388	371 4143 630	209 636 3.0e-62
Description		
pir: [LN:A69969] [AC:A69969] [PN:con [OR:Bacillus subtilis] [DB:pir2] >gp [AC:Z99116:AL009126] [GN:yqzB] [FN:u [DE:Bacillus subtilis complete genom [NT:similar to hypothetical proteins >gp: [GI:e1183755:g2634971] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-b 14 of 21): from 2599451to 2812870.] [RE:5423] [DI:complement]	[GI:e1185791:g263499] known] [OR:Bacillus (section 13 of 21) [LE:208975] [RE:209] 14] [AC:Z99117:AL009] t1] [DE:Bacillus sul	57] [LN:BSUB0013] subtilis] [DB:genpept-bct1] : from 2395261to 2613730.] 9613] [DI:complement] 9126] [GN:yqzB] [FN:unknown] btilis complete genome (section

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000982_26209425_c1_1022	372	4144	1068	355	1672	1.9e-172
Description sp:[LN:RPSD_STAAU] [AC:P26766] [GN:FPDLYMERASE SIGMA FACTOR RPOD] [SP:P2 [PN:transcription initiation factor factor sigma 43: transcription initiation factor sigma region 1 hor page:[GI:g153069] [LN:STAPLAC] [AC:M6 [OR:Staphylococcus aureus] [SR:Staph [DB:genpept-bct1] [DE:S.aureus sigma [RE:1926] [DI:direct]	26766] [D sigma pl iation fa mology] [53177] [P nylococcu	DB:swis .aC] [0 .ctor s .OR:Sta .DN:sign .s aure	ssprot] SN:plaC sigma k aphyloc na fact eus (st	>pir: [CL atF ho occus or] [G rain S	[LN:S34 c:transo mology aureus] EN:plaC]	A442] [AC:S34442] cription initiation transcription [DB:pir2] [NA]
ORF Name	NTID	AAID	<u>NT</u> Lengthl	<u>AA</u> Length	score	probability
A17503000982_26212501_c3_1394	373	4145	852	283	797 2	2.6e-79
Description			, L	-		
gp:[GI:g5019735] [LN:AF104349] [AC:R [GN:zurM] [OR:Listeria monocytogeness zinc-like uptake operon, complete se	s] [DB:ge	npept-	bct2]	[DE:Li	steria	monocytogenes
ORF Name	NTID	AAID	<u>NT</u> Lengthl	<u>AA</u> Length	score	probability
AI7503000982_26225463_c1_1070	374	4146	741	246	821 7	.4e-82
Description sp:[LN:RESD_BACSU] [AC:P35163] [GN:FREGULATORY PROTEIN RESD] [SP:P35163]] [PN:two-component response regulator >gp:[GI:g410141] [LN:BACDIA] [AC:L09(strain 168, sub_species Marburg) DN serA region.] [NT:ORFX17] [LE:21706] [LN:BSUB0013] [AC:Z99116:AL009126] [FN:activation role in global regulation global regulation global regulation complement] [DI:complement]	[DB:swi cor invol homology 228] [OR VA] [DB:g [RE:224 [PN:two-cation of ilis comp	ssprot ved in [OR: :Bacil enpept 28] [I compone aerobi	c] >pir n aerob Bacill lus su c-bctl] DI:direc ent resp cc] [OR	:[LN:G ic and us sub btilis [DE:B ct] >g ponse :Bacil (secti	69691] anaer tilis]] [SR:F acillus p:[GI:e regulat lus suk on 13 c	[AC:G69691:S45559 resD] [GN:resD] [DB:pir2] Bacillus subtilis s subtilis spoVA to e1185581:g2634747] cor] [GN:resD] otilis] of 21): from
ORF Name A17503000982_26251577 c1 994			NT LengthI	Length	score	probability
Description	375	4147	354	117	372 2	.8e-34
sp:[LN:YQEL_BACSU] [AC:P54457] [GN:YKD PROTEIN IN AROD-COMER INTERGENIC [AC:H69951] [PN:ybeB protein homoloprotein] [OR:Bacillus subtilis] [DB:[AC:D84432:D82370] [PN:YqeL] [OR:Baciltrain:JH642(trpC2 PheA1)) DNA] [DEregion containing skin element.] [LE>gp:[GI:e1183792:g2635008] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-b	REGION] og yqeL] pir2] >g cillus su 3:genpept 1:76417]	[SP:P5 [GN:yqp:[GI: btilis -bct1] [RE:76 :29911	4457] [eL] [0 d101312] [SR:1 [DE:Ba 773] [I 7:AL009	[DB:sw. CL:Escl 28:g13 Bacillu acillu: DI:dire 9126]	issprot herichi 03793] us subti s subti ect] [GN:yqe	:] >pir:[LN:H69951] .a coli ybeB [LN:BACJH642] :ilis .lis DNA, 283 Kb

14 of 21): from 2599451to 2812870.] [NT:similar to hypothetical proteins] [SP:P54457]

[LE:42657] [RE:43013] [DI:complement]

NT ORF Name NTID score probability LengthLength AI7503000982 26259657 cl 1051 4148 1425 1155 3.0e-117 Description sp:[LN:DLD2 BACSU] [AC:P54533] [GN:BFMBC] [OR:BACILLUS SUBTILIS] [EC:1.8.1.4] [DE:DEHYDROGENASE) (LPD-VAL)] [SP:P54533] [DB:swissprot] NT AΑ ORF Name NTID AAID <u>score</u> probability LengthLength A17503000982_26306713 c2 1332 377 4149 912 303 580 2.6e-56 Description pir:[LN:S72642] [AC:S72642] [PN:probable ABC-type transport protein xynB] [GN:xynB] [OR:Thermoanaerobacterium thermosulfurigenes] [DB:pir2] >gp:[GI:q1255237] [LN:TTU50952] [AC:U50952] [PN:XynB] [GN:xynB] [OR:Thermoanaerobacterium thermosulfurigenes] [DB:genpept-bct1] [DE:Thermoanaerobacterium thermosulfurigenes endoxylanase precursor(XynA) and membrane component of an ABC transporter (XynB) genes, complete cds and XynC (xynC) gene, partial cds.] [NT:Description: membrane component of an ABC] [LE:454] [RE:1359] [DI:direct] NT AA ORF Name NTID AAID score probability Length Length A17503000982 26368950 c2 1264 378 4150 1380 459 4.7e-76 Description sp:[LN:RECQ BACSU] [AC:P50729] [GN:RECQ] [OR:BACILLUS SUBTILIS] [EC:3.6.1.-] [DE:ATP-DEPENDENT DNA HELICASE RECQ,] [SP:P50729] [DB:swissprot] >pir:[LN:A69691] [AC:A69691] [PN:ATP-dependent DNA helicase homolog:ATP-dependent DNA helicase recQ] [GN:recQ] [CL:DEAD/H box helicase homology] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1183747:g2634720] [LN:BSUB0012] [AC:Z99115:AL009126] [PN:ATP-dependent DNA helicase] [GN:recQ] [OR:Bacillus subtilis] [DB:genpept-bct1] [EC:3.6.1.-] [DE:Bacillus subtilis complete genome (section 12 of 21): from 2195541to 2409220.] [SP:P50729] [LE:210616] [RE:212106] [DI:complement] >gp:[GI:e1185571:q2634737] [LN:BSUB0013] [AC:Z99116:AL009126] [PN:ATP-dependent DNA helicase] [GN:recQ] [OR:Bacillus subtilis] [DB:genpept-bct1] [EC:3.6.1.-] [DE:Bacillus subtilis complete genome (section 13 of

21): from 2395261to 2613730.] [SP:P50729] [LE:10896] [RE:12386] [DI:complement] >gp:[GI:g1146200] [LN:BACSERA] [AC:L47648] [PN:DNA or RNA helicase, DNA-dependent ATPase] [GN:recS] [FN:DNA repair and homologous recombination] [OR:Bacillus subtilis]

[DB:genpept-bct2] [DE:Bacillus subtilis phosphoglycerate dehydrogenase (serA),

monophosphatekinase (cmk), ypfD, ypgA, yphA, yphB, yphC, NAD+

ypaA, ferredoxin (fer), ypbB, recS, ypbD, ypbE, ypbF, ypbG, ypbH, glutamate dehydrogenase (ypcA), ypdA, ypdB, ypdC, spore cortexlytic enzyme (sleB), ypeB, ypfA, ypfB, cytidine

dependentglycerol-3-phosphate dehydrogenase (glyc), yphE and yphF genes,complete cds.] [NT:similar to Escherichia coli recQ gene product;] [LE:4291] [RE:5781] [DI:direct]

NT AA ORF Name AAID NTID score probability LengthLength

AI7503000982 26384682 cl 1092

[LE:170048] [RE:171319] [DI:complement]

415

2.6e-54

Description

sp:[LN:YPIA BACSU] [AC:P54389] [GN:YPIA] [OR:BACILLUS SUBTILIS] [DE:HYPOTHETICAL 48.3 KD PROTEIN IN QCRA-AROE INTERGENIC REGION] [SP:P54389] [DB:swissprot] >pir:[LN:E69936] [AC:E69936] [PN:conserved hypothetical protein ypiA] [GN:ypiA] [CL:tetratricopeptide repeat homology] [OR:Bacillus subtilis] [DB:pir2] >qp:[GI:g1146224] [LN:BACYPIA] [AC:L47709] [GN:ypiA] [FN:hypothetical] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis (clone YAC15-6B) ypiABF genes, qcrABC genes, ypjABCDEFGHI genes, birA gene, panBCD genes, dinG gene, ypmB gene, aspB gene, asnS gene, dnaD gene, nth gene and ypoC gene, completecds's.] [NT:putative] [LE:348] [RE:1619] [DI:direct] >gp:[GI:e1183704:g2634677] [LN:BSUB0012] [AC:Z99115:AL009126] [GN:ypiA] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 12 of 21): from 2195541to 2409220.] [NT:similar to hypothetical proteins] [SP:P54389]

NT AA ORF Name NTID <u>sco</u>re AAID probability LengthLength AI7503000982 26597186 cl 1056 380 444 147

Description

sp:[LN:YQIW BACSU] [AC:P54534] [GN:YOIW] [OR:BACILLUS SUBTILIS] [DE:HYPOTHETICAL 16.2 KD PROTEIN IN BMRU-ANSR INTERGENIC REGION] [SP:P54534] [DB:swissprot] >pir:[LN:E69962] [AC:E69962] [PN:conserved hypothetical protein yqiW] [GN:yqiW] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:d1013283:g1303948] [LN:BACJH642] [AC:D84432:D82370] [PN:YqiW] [OR:Bacillus subtilis] [SR:Bacillus subtilis (strain:JH642(trpC2 PheA1)) DNA] [DB:genpept-bct1] [DE:Bacillus subtilis DNA, 283 Kb region containing skin element.] [LE:226148] [RE:226585] [DI:direct] >gp:[GI:e1185667:g2634833] [LN:BSUB0013] [AC:Z99116:AL009126] [GN:yqiW] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 13 of 21): from 2395261to 2613730.] [NT:similar to hypothetical proteins from B. subtilis] [LE:97034] [RE:97471] [DI:complement]

NTAΑ ORF Name score NTID AAID probability LengthLength A17503000982_26600137_f2_493 381 4153 8.9e-15 531

Description

sp:[LN:YPUF BACSU] [AC:P17617] [GN:YPUF] [OR:BACILLUS SUBTILIS] [DE:HYPOTHETICAL 21.0 KD PROTEIN IN RIBT-DACB INTERGENIC REGION (ORFX6)] [SP:P17617] [DB:swissprot] >pir:[LN:S45548] [AC:S45548:E69942] [PN:hypothetical protein ypuF] [GN:ypuF] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:g410130] [LN:BACDIA] [AC:L09228] [OR:Bacillus subtilis] [SR:Bacillus subtilis (strain 168, sub species Marburg) DNA] [DB:genpept-bct1] [DE:Bacillus subtilis spoVA to serA region.] [NT:ORFX6] [LE:12218] [RE:12742] [DI:complement] >qp:[GI:q580916] [LN:BSRIB] [AC:X51510] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:B.subtilis riboflavin biosynthesis operon ribG, ribB, ribA, ribH, and ribT genes.] [NT:ORF Y (AA 1-174)] [SP:P17617] [LE:5164] [RE:5688] [DI:complement] >gp:[GI:e1185592:g2634758] [LN:BSUB0013] [AC:Z99116:AL009126] [GN:ypuF] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 13 of 21): from 2395261to 2613730.] [SP:P17617] [LE:30841] [RE:31365] [DI:direct]

ORF Name probability LengthLength AI7503000982 26752312 c3 1396 382 171 56 4.0e-19 sp:[LN:RL33 BACST] [AC:P23375] [GN:RPMG] [OR:BACILLUS STEAROTHERMOPHILUS] [DE:50S RIBOSOMAL PROTEIN L33] [SP:P23375] [DB:swissprot] >pir:[LN:B48396] [AC:B48396] [PN:ribosomal protein L33] [CL:Escherichia coli ribosomal protein L33] [OR:Bacillus stearothermophilus] [DB:pir2] NTAA ORF Name NTID AAID Length Length score probability AI7503000982 26754662 c3 1454 383 2.5e-49 Description sp:[LN:KCY BACSU] [AC:P38493] [GN:CMK:JOFC] [OR:BACILLUS SUBTILIS] [EC:2.7.4.14] [DE:(CMP KINASE)] [SP:P38493] [DB:swissprot] >pir:[LN:F69601] [AC:F69601] [PN:cytidylate kinase cmk] [GN:cmk] [CL:cytidylate kinase cmk] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:g533105] [LN:BSU11687] [AC:U11687] [PN:unknown] [GN:jofC] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis 168 jofA, jofB, MssA homolog (jofC) and ribosomalprotein S1 homolog (jofD) genes, complete cds, and joeB gene, partial cds.] [NT:similar to the Escherichia coli mssA gene product] [LE:1744] [RE:2418] [DI:direct] >gp:[GI:e1183734:g2634707] [LN:BSUB0012] [AC:Z99115:AL009126] [PN:cytidylate kinase] [GN:cmk] [OR:Bacillus subtilis] [DB:genpept-bct1] [EC:2.7.4.14] [DE:Bacillus subtilis complete genome (section 12 of 21): from 2195541to 2409220.] [NT:alternate gene name: jofC, ypfC] [SP:P38493] [LE:199742] [RE:200416] [DI:complement] >gp:[GI:e1185558:g2634724] [LN:BSUB0013] [AC:Z99116:AL009126] [PN:cytidylate kinase] [GN:cmk] [OR:Bacillus subtilis] [DB:genpept-bct1] [EC:2.7.4.14] [DE:Bacillus subtilis complete genome (section 13 of 21): from 2395261to 2613730.] [NT:alternate gene name: jofC, ypfC] [SP:P38493] [LE:22] [RE:696] [DI:complement] >gp:[GI:g1146214] [LN:BACSERA] [AC:L47648] [PN:cytidine monophosphate kinase] [GN:cmk] [FN:cytidine diphosphate biosynthesis] [OR:Bacillus subtilis] [DB:genpept-bct2] [DE:Bacillus subtilis phosphoglycerate dehydrogenase (serA), ypaA, ferredoxin (fer), ypbB, recS, ypbD, ypbE, ypbF, ypbG, ypbH, glutamate dehydrogenase (ypcA), ypdA, ypdB, ypdC, spore cortexlytic enzyme (sleB), ypeB, ypfA, ypfB, cytidine monophosphatekinase (cmk), ypfD, ypgA, yphA, yphB, yphC, NAD+ dependentglycerol-3-phosphate dehydrogenase (glyc), yphE and yphF genes, complete cds.] [NT:similar to Escherichia coli smba supress; putative] [LE:15981] [RE:16655] [DI:direct] NT AA ORF Name NTID AAID probability LengthLength AI7503000982 26754750 c2 1172 384 4156 1164 387 L.1e-89 Description

pir:[LN:E69981] [AC:E69981] [PN:NifS protein homolog homolog yrvO] [GN:yrvO] [CL:nitrogen fixation protein nifS] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1183981:g2635197] [LN:BSUB0014] [AC:Z99117:AL009126] [GN:yrvO] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 14 of 21): from 2599451to 2812870.] [NT:similar to NifS protein homolog] [LE:210475] [RE:211515] [DI:complement] >qp:[GI:e1183999:q2635215] [LN:BSUB0015] [AC:Z99118:AL009126] [GN:yrvO] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 15 of 21): from 2795131to 3013540.] [NT:similar to NifS protein homolog] [LE:14795] [RE:15835] [DI:complement]

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000982_26772135_c1_1000	385	4157		331	555 1	1e-53
Description sp:[LN:YQEN_BACSU] [AC:P54459] [GN:Y KD PROTEIN IN COMEC-RPST INTERGENIC [AC:B69952] [PN:conserved hypotheti [DB:pir2] >gp:[GI:d1013134:g1303799] [OR:Bacillus subtilis] [SR:Bacillus [DB:genpept-bct1] [DE:Bacillus subti [LE:82475] [RE:83518] [DI:direct] >g [AC:Z99117:AL009126] [GN:yqeN] [FN:u [DE:Bacillus subtilis complete genom [NT:similar to hypothetical proteins	REGION) cal pro [LN:BA subtili lis DNA pp:[GI:e unknown] ne (sect	[SP:F tein y CJH642 s (str , 283 118378 [OR:E	P54459] PqeN] [G Pain:JH6 Kb regi P6:g2635 Pacillus Pof 21)	[DB:sw. N:yqeN 84432:1 42(trpe on con- 002] [1 subtil : from	issprot [OR: 082370] C2 Phea taining LN:BSUE lis] [I	c] >pir:[LN:B69952] :Bacillus subtilis] [PN:YqeN] A1)) DNA] g skin element.] B0014] DB:genpept-bct1] Sito 2812870.]
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
AI7503000982_26774062_c2_1268	386	4158	1332	443	1670 8	3.0e-172
Description sp:[LN:YPHC_BACSU] [AC:P50743] [GN:Y [DB:swissprot] >pir:[LN:A69936] [AC: [GN:yphC] [CL:Mycobacterium leprae factor Tu homology] [OR:Bacillus sub [LN:BSUB0012] [AC:Z99115:AL009126] [[DB:genpept-bct1] [DE:Bacillus subti 2195541to 2409220.] [NT:similar to b [DI:complement] >gp:[GI:g1146219] [I subtilis] [DB:genpept-bct2] [DE:Baci (serA), ypaA, ferredoxin (fer), ypbB, dehydrogenase (ypcA), ypdA, ypdB, yp ypfB, cytidine monophosphatekinase (dependentglycerol-3-phosphate dehydr [NT:similar to Escherichia coli GTP-	A69936 probabl tilis] GN:yphC lis com ypothet N:BACSE llus su recS, dC, spo cmk), y ogenase] [PN: e GTP- [DB:pi] [FN: plete ical p RA] [A btilis ypbD, re cor pfD, y (glyc	conserv binding r2] >gp unknown genome roteins C:L4764 phosph ypbE, y texlyti pgA, yp), yphE	ed hypo protes: [GI:e] [OR:I (section] [LE:1 8] [GN:1 oglyces pbF, yr c enzyces hA, yph and yr	othetic in:tran 1183729 Bacillu on 12 c 193903] :yphC] rate de pbG, yp me (sle nB, yph phF gen	cal protein yphC] islation elongation 0:g2634702] is subtilis] of 21): from [RE:195213] [OR:Bacillus chydrogenase obH,glutamate eB), ypeB, ypfA, ic, NAD+ ies,complete cds.]
ORF Name A17503000982_2756288_c2_1257 Description NO-HIT	<u>NTID</u>	<u>AAID</u> 4159	NT Lengthi	AA Length ⁵	score	probability
ORF Name A17503000983_10000128_f2_399	NTID 388	<u>AAID</u> 4160	NT Length1	AA Length	score	probability

Description NO-HIT

DRF Name	NTID	AAID	NT LengthL	AA ength score	probabi	lity
AI7503000983_10195942_c3_1056	389	4161		211 253	1.2e-21	
Description Sp:[LN:TRPF_THEMA] [AC:Q56320] [GN:T [DE:N-(5'-PHOSPHORIBOSYL)ANTHRANILAT pir:[LN:S59048] [AC:S59048] [PN:ph [CL:phosphoribosylanthranilate iso [DB:pir2] >gp:[GI:g1054860] [LN:TMTR isomerase] [GN:trpF] [OR:Thermotoga trpC, trpF, trpB, and trpA genes.] [pgp:[GI:g4980631] [LN:AE001699] [AC: isomerase] [GN:TM0139] [OR:Thermotog maritima section 11 of 136 of the co GP:Q56320 GB:AE000512] [LE:7969] [RE	TE ISOME COSPHORIS COSPHOR	ERASE, bosyl trpF h [AC:X9 ma] [DB 220] [L 29:AE00 ima] [genome	(PRAI)] anthrani omology] 2729] [E :genpept E:1012] 0512] [E DB:genpe	[SP:Q56320 late isome [OR:Therm PN:phosphor :-bct1] [DE [RE:1629] PN:phosphor ept-bct2] [similar to	DB:swis rase trpF] otoga mari ibosyl ant :T.maritim [DI:direct ibosylanth DE:Thermot	sprot] [GN:trpF tima] hranilate a trpD,] ranilate oga
ORF Name A17503000983_10553766_£2_397	NTID	<u>AAID</u>		AA score	probabi	lity
Description NO-HIT		L	J			
ORF Name	NTID	AAID	<u>NT</u> LengthL	AA ength	probabi	lity
A17503000983_1058463_c3_978	391	4163	2121	706 2472	8.3e-257	
pir:[LN:S70691] [AC:S70691:C69680] alpha chain pnpA:polynucleotide phos nucleotidyltransferase alpha chain] pg:[GI:e1185260:g2634041] [LN:BSUB0 phosphorylase (PNPase)] [GN:pnpA] [F[OR:Bacillus subtilis] [DB:genpept-bgenome (section 9 of 21): from 15984 [SP:P50849] [LE:140354] [RE:142471] [AC:U29668] [PN:polynucleotide phosp [DB:genpept-bct2] [DE:Bacillus subticts, and polynucleotide phosphorylase [DI:direct]	phoryla [OR:Bac 009] [A N:neces ct1] [E 21to 18 [DI:dir horylas lis rib	see pnp. illus C:Z991 sary f C:2.7. 07200. ect] > e] [GN oosomal	A] [GN:psubtilis 12:AL009 or compe 7.8] [DE] [NT:al gp:[GI:g :pnpA] [protein	mpA] [CL: 2] [EC:2.7. 226] [PN:p 2: tence deve 4: Bacillus 4: ternate ge 2: 1184680] [OR: Bacillu 4: RpsO (rps	polyribonud 7.8] [DB:pi olynucleoti lopment] subtilis cone name: cone name: co LN:BSU29666 s subtilis O) gene, pa	cleotide ir2] ide omplete omR] 8] artial
ORF Name A17503000983_10657925_c3_1017 Description Oir: [LN:H69722] [AC:H69722:B25364] In the second of the secon	[EC:2.7 PN:homo [DB:ge of 21)	.1.39] serine npept-l : from	e kinase [DB:pir kinase] oct1] [E 3197001	,] [GN:thr 2] >gp:[GI [GN:thrB] C:2.7.1.39	1.4e-55 B] [CL:hor:e1184303:g [FN:threor] [DE:Baci]	moserine g2635721] nine llus
ORF Name A17503000983_10667002_f2_278 Description	<u>NTID</u>	<u>AAID</u> 4165	NT LengthL		probabil	.ity

ORF Name	NTID A	AAID NT AA score probability
A17503000983_1074090_c1_728	394 4	166 906 301 697 1.0e-68
Description	L L	
<pre>site-specific integrase/recombinase >gp:[GI:g535348] [LN:BSU13634] [AC:U</pre>	76] [DB:strase/recor XerC] [OR 13634] [PI lis JH642 plete cds 009] [AC: [OR:Bacile (section	wissprot] >pir:[LN:G69601] mbinase codV] [GN:codV] [CL:probable a:Bacillus subtilis] [DB:pir2] eN:CodV] [GN:codV] [OR:Bacillus subtilis] a dipeptide permease operon regulators, a.] [LE:293] [RE:1207] [DI:direct] Z99112:AL009126] allus subtilis] [DB:genpept-bct1] an 9 of 21): from 1598421to 1807200.]
ORF Name		AAID NT AA score probability
A17503000983_10975428_£2_340	395 4	879 292 80 0.013
[DE:CYTOSKELETON-LIKE BICAUDAL D PRO	TEIN] [SP [GN:BicD 1684] [OR embryo, cI bicaudalI	DNA to mRNA] [DB:genpept-inv1] D protein (BicD) mRNA,complete cds.]
ORF Name	NTID A	AAID NT AA score probability
ORF Name AI7503000983_1179775_f1_70		MID — gcore probability
AI7503000983_1179775_f1_70 Description pir:[LN:S78676] [AC:S78676:S78677] Saccharomyces cerevisiae] [DB:pir2] [AC:AJ011856] [OR:Mitochondrion Sacc	[PN:hypoth >gp:[GI:entharomyces cerevisiae	hetical protein Q0144] [OR:mitochondrion 1370708:g4160384] [LN:SCE011856]
AI7503000983_1179775_f1_70 Description pir:[LN:S78676] [AC:S78676:S78677] Saccharomyces cerevisiae] [DB:pir2] [AC:AJ011856] [OR:Mitochondrion Sacci	[PN:hypoth >gp:[GI:ei haromyces cerevisiae ect]	hetical protein Q0144] [OR:mitochondrion 1370708:g4160384] [LN:SCE011856] cerevisiae] [SR:baker's yeast]
AI7503000983_1179775_f1_70 Description pir:[LN:S78676] [AC:S78676:S78677] Saccharomyces cerevisiae] [DB:pir2] [AC:AJ011856] [OR:Mitochondrion Sacc. [DB:genpept-pln1] [DE:Saccharomyces Q0144] [LE:54109] [RE:54438] [DI:direction of the content of	[PN:hypoth >gp:[GI:ei haromyces cerevisiae ect]	LengthLength LengthLength LengthLength 132 43 42 0.029 hetical protein Q0144] [OR:mitochondrion 1370708:g4160384] [LN:SCE011856] cerevisiae] [SR:baker's yeast] e complete mitochondrial genome.] [NT:ORF
AI7503000983_1179775_f1_70 Description pir:[LN:S78676] [AC:S78676:S78677] Saccharomyces cerevisiae] [DB:pir2] [AC:AJ011856] [OR:Mitochondrion Sacci [DB:genpept-pln1] [DE:Saccharomyces Q0144] [LE:54109] [RE:54438] [DI:direct ORF Name AI7503000983_11817625_f1_65 Description NO-HIT ORF Name	[PN:hypoth >gp:[GI:e] haromyces cerevisiae ect] NTID A	LengthLength LengthLength LengthLength 168
AI7503000983_1179775_f1_70 Description pir:[LN:S78676] [AC:S78676:S78677] Saccharomyces cerevisiae] [DB:pir2] [AC:AJ011856] [OR:Mitochondrion Saccion [DB:genpept-pln1] [DE:Saccharomyces Q0144] [LE:54109] [RE:54438] [DI:director of the control of	[PN:hypoth >gp:[GI:e] haromyces cerevisiae ect] NTID A	LengthLength LengthLength 168 132 43 42 0.029 hetical protein Q0144] [OR:mitochondrion 1370708:g4160384] [LN:SCE011856] cerevisiae] [SR:baker's yeast] e complete mitochondrial genome.] [NT:ORF AID NT AA LengthLength 169 129 42

ORF Name A17503000983_11844802_c2_941	<u>NTID</u>	AAID <u>NT AA</u> LengthLength score probability 4171	
Description NO-HIT			
ORF Name	NTID	AAID NT AA LengthLength score probability	
A17503000983_11926627_f3_614	400	4172 171 56	
Description NO-HIT			
MO-HII			
ORF Name	NTID	$rac{ ext{NT}}{ ext{Length}}rac{ ext{AA}}{ ext{score}}$ probability	
AI7503000983_11930317_c3_974	401	4173 336 111 251 1.9e-21	
PROTEIN IN NUSA-INFB INTERGENIC REGI >pir:[LN:E36905] [AC:E36905:C69882:S ylxQ:hypothetical protein 2 (infB 5' [DB:pir2] >gp:[GI:g49318] [LN:BSORF1 [DB:genpept-bct1] [DE:B.subtilis inf [DI:direct] >gp:[GI:e1185253:g263403 [FN:unknown] [OR:Bacillus subtilis]	ON (ORF 31993] region T7A] [A B-nusA 4] [LN: [DB:gen 21to 18	[PN:ribosomal protein L7AE family hom [GN:ylxQ] [OR:Bacillus subtilis] 218631] [GN:ORF4] [OR:Bacillus subticeron.] [SP:P32729] [LE:2367] [RE:266 [SUB0009] [AC:Z99112:AL009126] [GN:ylxept-bct1] [DE:Bacillus subtilis comploated of the comploation of the complex com	olog lis] 9] Q]
ORF Name	NTID	AAID NT AA score probability	
ORF Name AI7503000983_1212785_f1_108	<u>NTID</u>	AAID <u>NT AA</u> LengthLength score probability 4174 213 70 74 0.048	
		LengthLength probability	
Description pir: [LN:G70047] [AC:G70047] [PN:con [OR:Bacillus subtilis] [DB:pir2] >gp [AC:Z99120:AL009126] [GN:yvrM] [FN:u [DE:Bacillus subtilis complete genom [NT:similar to hypothetical proteins >gp: [GI:e1186014:g2635839] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-b	served : :[GI:e1:mknown] ne (sect:] [LE:2:018] [A:oct1] [Di	LengthLength score probability 4174 213 70 74 0.048 Prothetical protein yvrM] [GN:yvrM] 84405:g2635823] [LN:BSUB0017] [OR:Bacillus subtilis] [DB:genpept-boon 17 of 21): from 3197001to 3414420.	wn] ection
Description pir: [LN:G70047] [AC:G70047] [PN:conform [OR:Bacillus subtilis] [DB:pir2] > gp [AC:Z99120:AL009126] [GN:yvrM] [FN:ufform [NT:similar to hypothetical proteins > gp: [GI:e1186014:g2635839] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-b18 of 21): from 3399551to 3609060.]	served : :[GI:e1:mknown] ne (sect:] [LE:2:018] [A:oct1] [Di	LengthLength LengthLength LengthLength A174 [A174 [A1	wn] ection
Description pir: [LN:G70047] [AC:G70047] [PN:con [OR:Bacillus subtilis] [DB:pir2] >gp [AC:Z99120:AL009126] [GN:yvrM] [FN:u [DE:Bacillus subtilis complete genom [NT:similar to hypothetical proteins >gp: [GI:e1186014:g2635839] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-b 18 of 21): from 3399551to 3609060.] [RE:12401] [DI:complement]	served : [GI:el : mknown] [LE:2: 018] [A: ct1] [D: [NT:sim.	LengthLength LengthLength LengthLength A174 LengthLength For a content of the content of th	wn] ection
Description pir: [LN:G70047] [AC:G70047] [PN:con [OR:Bacillus subtilis] [DB:pir2] >gp [AC:Z99120:AL009126] [GN:yvrM] [FN:u [DE:Bacillus subtilis complete genom [NT:similar to hypothetical proteins >gp: [GI:e1186014:g2635839] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-b 18 of 21): from 3399551to 3609060.] [RE:12401] [DI:complement] ORF Name	served :: [GI:e1: mknown] [He (sect: [] [LE:2: [] [All [] [NT:sim. [] [NT:sim.	LengthLength LengthLength LengthLength From [GN:yvrM] [GN:yvrM] [GR:Bacillus subtilis] [DB:genpept-boon 17 of 21): from 3197001to 3414420. [A190] [RE:214951] [DI:complement] [EXEMPLIE SUBTILITY SERVICE SUBTILITY SUBTILIT	wn] ection
Description pir: [LN:G70047] [AC:G70047] [PN:com [OR:Bacillus subtilis] [DB:pir2] >gp [AC:Z99120:AL009126] [GN:yvrM] [FN:u [DE:Bacillus subtilis complete genom [NT:similar to hypothetical proteins >gp: [GI:e1186014:g2635839] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-b 18 of 21): from 3399551to 3609060.] [RE:12401] [DI:complement] ORF Name A17503000983_12694082_c2_889	served :: [GI:e1: mknown] [He (sect: [] [LE:2: [] [All [] [NT:sim. [] [NT:sim.	LengthLength LengthLength LengthLength From [GN:yvrM] [GN:yvrM] [GR:Bacillus subtilis] [DB:genpept-boon 17 of 21): from 3197001to 3414420. [A190] [RE:214951] [DI:complement] [EXEMPLIE SUBTILITY SERVICE SUBTILITY SUBTILIT	wn] ection
Description pir: [LN:G70047] [AC:G70047] [PN:con [OR:Bacillus subtilis] [DB:pir2] >gp [AC:Z99120:AL009126] [GN:yvrM] [FN:u [DE:Bacillus subtilis complete genom [NT:similar to hypothetical proteins >gp: [GI:e1186014:g2635839] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-b 18 of 21): from 3399551to 3609060.] [RE:12401] [DI:complement] ORF Name AT7503000983_12694082_c2_889 Description	served :: [GI:e1: mknown] [He (sect: [] [LE:2: [] [All [] [NT:sim. [] [NT:sim.	LengthLength LengthLength LengthLength From [GN:yvrM] [GN:yvrM] [GR:Bacillus subtilis] [DB:genpept-boon 17 of 21): from 3197001to 3414420. [A190] [RE:214951] [DI:complement] [EXEMPLIE SUBTILITY SERVICE SUBTILITY SUBTILIT	wn] ection

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000983_13087513_c2_907	405	4177	3033	1010	417	4.1e-65
Description						
pir:[LN:D71302] [AC:D71302] [PN:problems of the complete genome.] [NT:s:[LE:12418] [RE:15561] [DI:direct]	um] [SR AE00123 [DB:gen	:, sypl 7:AE000 pept-bo	nilis s 0520] [ct2] [D	piroch PN:exc E:Trep	ete] nuclea onema	[DB:pir2] ase (sbcC)] pallidum section 53
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
AI7503000983_13179692_c1_724	406	4178	132	43		
Description					•	
NO-HIT						
ORF Name	NTID		NT Length		score	probability
A17503000983_1360958_c2_905 Description	407	4179	270	89	109	2.1e-06
pir:[LN:C69891] [AC:C69891:S57406] [OR:Bacillus subtilis] [DB:pir2] >gp [PN:YneJ] [GN:yneJ] [OR:Bacillus subtilis] [PN:YneJ] [GN:yneJ] [N:G8:G8:G8:G8:G8:G8:G8:G8:G8:G8:G8:G8:G8:	:[GI:e2 tilis] T:ident direct] nknown] e (sect	49647:g [DB:ger ical to >gp:[0 [OR:Ba ion 10	g140545 npept-b yoxI GI:e118 acillus of 21)	1] [LN ct1] [(from 3454:g subti : from	[:BC170 [DE:B.s acc. r 263417 [lis] 17812	DDEGR] [AC:Z73234] Subtilis DNA (26.2 no. X87845)] 79] [LN:BSUB0010] [DB:genpept-bct1] 201to 2014980.]
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000983_13678135_£3_575	408	4180		52		0.00093
Description pir:[LN:B71622] [AC:B71622] [PN:meta [GN:PFB0140w] [OR:Plasmodium falcipa						
[AC:AE001375:AE001362] [PN:metal bind [OR:Plasmodium falciparum] [SR:malar: [DE:Plasmodium falciparum chromosome [NT:identified by sequence similarity	ding pro ia para 2, sec	otein site P. tion 12	(DHHC d falci of 73	omain) parum] of th] [GN: [DB:g ecomp]	PFB0140w] genpept-inv2] .ete sequence.]
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000983_13678462_f1_15	409	4181	564	187	303	5.8e-27
Description					·	
sp:[LN:SPHX_SYNP7] [AC:P39665] [GN:SINIDULANS R2] [DE:SPHX PROTEIN PRECURS [AC:S54402] [PN:SphX protein] [CL:sp>gp:[GI:d1005689:g496319] [LN:SYOSPHX [OR:Synechococcus sp.] [SR:Synechococcus sp.] [SR:Synechococcus sp.] [SR:Synechococcus sp.] [RE:1330] [DI:direct]	SOR] [SI phX prof [AC:1 ccus sp	P:P3966 tein] D26161] . (stra	55] [DB [OR:Syn [PN:S iin:PCC	:swiss echoco phX] [7942)	prot] ccus s GN:sph DNA] [>pir:[LN:S54402] sp.] [DB:pir2] sX] DB:genpept-bct1]
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000983_13691280_t1_42	410	4182		44		
Description						
NO-HIT						

			NT(T) 7. 7.
ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000983_13845300_£2_231	411	4183	309 102 79 0.0065
Description		L	
] [PN:protein VI (trpA 3'-region)] [LN:BATRYOPEA] [AC:Z19055] [PN:ORF 6	[DB:s] [OR:Buch: [OR:Buch: [NT:ho	wisspr nera a uchner mologo	rot] >pir:[LN:F49897] [AC:F49897:S36433 aphidicola] [DB:pir2] >gp:[GI:g396663]
ORF Name	NTID	AAID	NT AA LengthLength score probability
AI7503000983_13875216_c3_1046	412	4184	1023 340 895 1.1e-89
Description		-	
gp:[GI:e1181777:g2632149] [LN:SCY140 [OR:Staphylococcus carnosus] [DB:gen [LE:213] [RE:1076] [DI:direct]			29] [PN:antiterminator] [GN:glcT] DE:Staphylococcus carnosus glcT gene.]
ORF Name	NTID	AAID	NT AA LengthLength score probability
AY7503000983_14251643_£3_547	413	4185	309 102 143 5.2e-10
Description			
pir:[LN:H69891] [AC:H69891] [PN:hypsubtilis] [DB:pir2] >gp:[GI:e249654:[GN:yneR] [OR:Bacillus subtilis] [DB fragment; 170 degree region).] [LE:1>gp:[GI:e1183464:g2634189] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-b of 21): from 1781201to 2014980.]	g1405458 genpept 3221] [1 010] [Adoct1] [D1	8] [LN t-bct1 RE:135 C:Z991 E:Baci	N:BC170DEGR] [AC:Z73234] [PN:YneR] L] [DE:B.subtilis DNA (26.2 kb 508] [DI:complement] L13:AL009126] [GN:yneR] [FN:unknown] Llus subtilis complete genome (section
ORF Name	NTID	AAID	NT AA score probability
A17503000983_14457876_f3_521	414	4186	135 44 194 2.1e-15
Description			
gp:[GI:g1022726] [LN:SHU35635] [AC:U haemolyticus] [SR:Staphylococcus hae [DE:Staphylococcus haemolyticus IS12 [LE:1101] [RE:1922] [DI:complement]	molytic	us str	ain=Y176] [DB:genpept-bct1]
ORF Name	NTID	AAID	NT AA score probability
AT7503000983_14471938_c3_1027	415	4187	183 60 231 2.5e-19
Description	L		
sp:[LN:RL33_BACST] [AC:P23375] [GN:R RIBOSOMAL PROTEIN L33] [SP:P23375] [[PN:ribosomal protein L33] [CL:Esche	DB:swiss	sprot]	>pir:[LN:B48396] [AC:B48396]

stearothermophilus] [DB:pir2]

ORF Name	NTID	AAID	<u>NT</u> LengthI	<u>AA</u> Length	score	probability	
AI7503000983 1461588 f1 16	416	4188		313	712	2.6e-70	
Description			الــــال		L		
pir:[LN:D69419] [AC:D69419] [PN:pho homolog] [CL:phoW protein] [OR:Archa [LN:AE001010] [AC:AE001010:AE000782] [GN:AF1357] [OR:Archaeoglobus fulgion section 97 of 172 of the complete gen PID:1303855] [LE:5294] [RE:6187] [DI	eoglobu [PN:ph dus] [DB enome.]	s fulg osphat :genpe [NT:si	idus] [I e ABC tı pt-bct2]	DB:pir ranspo [DE:	2] >g] rter, Archa	p:[GI:g2649218] permease protein] eoglobus fulgidus	
ORF Name	NTID	AAID	<u>NT</u> LengthI	<u>AA</u>	score	probability	_
A17503000983_14634450_f1_1	417	4189		67			
Description			JL				
NO-HIT							
ORF Name	NTID	AAID	<u>NT</u> Lengthl	AA ength	score	probability	-
AI7503000983_14723387_c2_904	418	4190		93	250	2.4e-21	
Description							
PROTEIN IN TTK-CCDA INTERGENIC REGION] [SP:P45708] [DB:swissprot] >pir:[LN:S57403] [AC:S57403:A69891] [PN:hypothetical protein yneF] [GN:yneF] [CL:conserved hypothetical protein yneF] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e249644:g1405448] [LN:BC170DEGR] [AC:Z73234] [PN:YneF] [GN:yneF] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:B.subtilis DNA (26.2 kb fragment; 170 degree region).] [NT:identical to yoxG (from acc. no. X87845)] [SP:P45708] [LE:4225] [RE:4443] [DI:direct] >gp:[GI:g870924] [LN:BSTKTCCDA] [AC:X87845] [GN:orf] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:B.subtilis ORF120, ORF160, tkt and ccdA genes.] [SP:P45708] [LE:1011] [RE:1229] [DI:direct] >gp:[GI:e1183450:g2634175] [LN:BSUB0010] [AC:Z99113:AL009126] [GN:yneF] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 10 of 21): from 1781201to 2014980.] [NT:alternate gene name: yoxG] [SP:P45708] [LE:140604] [RE:140822] [DI:direct]							
ORF Name	NTID	AAID	<u>NT</u> LengthL			probability	-
A17503000983_14850082_f1_18	419	4191	933	310	880	4.2e-88	
Description sp:[LN:YQGK_BACSU] [AC:P46342] [GN:Y (ORF75)] [SP:P46342] [DB:swissprot] transporter (ATP-binding pro) homolo malK:ATP-binding cassette homology] >gp:[GI:d1013193:g1303858] [LN:BACJH subtilis] [SR:Bacillus subtilis (str [DE:Bacillus subtilis DNA, 283 Kb re [RE:142395] [DI:direct] >gp:[GI:d101 [OR:Bacillus subtilis] [SR:Bacillus [DB:genpept-bct1] [DE:Bacillus subti geneproducts.] [NT:Similality to pho [RE:4564] [DI:direct] >gp:[GI:e11857]	>pir:[Lipg yqgK] [OR:Bac [642] [Ac ain:JH6 gion co 0228:g9 subtili lis DNA	N:E699 [GN:yeillus: C:D844: 42(trpentaining) 03307] s (stranspect	56] [AC: qgK] [C subtilis 32:D8237 C2 PheA1 ng skin [LN:BAC ain:JH64 cmologue ort prot	E69956 L:inne [] [DB: [] [PI]) DNI elemen [PST] [] (try es of t	5] [Fer mem :pir2] V:YqgK A] [DE nt.] [[AC:D5 pC2 ph the E.	PN:phosphate ABC abrane protein [C] [OR:Bacillus B:genpept-bct1] [LE:141613] [S8414] [PN:ORF75] [BeA1)) DNA] [Coli pst [Df] [LE:3782]	

[GN:yqgK] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 13 of 21): from 2395261to 2613730.] [NT:alternate gene name: yzmF; similar to phosphate ABC] [SP:P46342] [LE:181226] [RE:182008] [DI:complement]

ORF Name	NTID	AAID	NT AA LengthLength score probability	
A17503000983_14876553_c2_919	420	4192	996 331 570 2.9e-55	
Description gp:[GI:g1147744] [LN:EHU42211] [AC:U [FN:involved in the regulation of pe [DE:Enterococcus hirae PBP 5 synthes [RE:1627] [DI:direct]	nicilli	n] [OR	R:Enterococcus hirae] [DB:genpept-bct2]	
ORF Name	NTID	AAID	NT AA score probability	
AY7503000983_14901512_c3_1012	421	4193		
Description sp:[LN:NUC_STAIN] [AC:P43269] [GN:NUC_EC:3.1.31.1] [DE:(STAPHYLOCOCCAL NUT) >pir:[LN:S26079] [AC:S26079] [PN:th [OR:Staphylococcus intermedius] [DB: [PN:thermonuclease] [GN:nuc] [OR:Sta [DE:S.intermedius nuc gene for therm [DI:direct]	CLEASE) ermonuc pir2] >] [SP: lease] gp:[GI ccus i	:P43269] [DB:swissprot]] [CL:micrococcal nuclease] I:g47146] [LN:SINUC8] [AC:X67678] intermedius] [DB:genpept-bct1]	
ORF Name	NTID	AAID	NT AA LengthLength score probability	-
A17503000983_15084826_c3_1034	422	4194		
Description sp:[LN:CCDC_BACSU] [AC:P45710] [GN:C [SP:P45710] [DB:swissprot] >gp:[GI:g [OR:Bacillus subtilis] [DB:genpept-b genes.] [SP:P45710] [LE:2942] [RE:34	870927] ct1] [D	[LN:B E:B.su	BSTKTCCDA] [AC:X87845] [GN:ORF160] ubtilis ORF120, ORF160, tkt and ccdA	
ORF Name	NTID	AAID	NT AA LengthLength score probability	_
A17503000983_15757712_f1_23	423	4195	732 243 686 1.5e-67	
Description gp:[GI:g3800828] [LN:AF076684] [AC:A ATPase domain] [GN:opp-2F] [OR:Staph [DE:Staphylococcus aureus oligopepti (opp-2B), oligopeptide transporter p oligopeptide transporterputative ATP transporterputative ATPase domain (o [DI:direct]	ylococc de tran utative ase dom	us auro sporte: membra ain (o	reus] [DB:genpept-bct2] er putative membranepermease domain ane permease domain (opp-2C), opp-2D), and oligopeptide	
ORF Name	NTID	AAID	NT AA LengthLength score probability	_
A17503000983_15803510_c2_854 Description	424	4196	132 43	
NO-HIT		*******		_
ORF Name			NT AA	
	NTID	AAID	LengthLength score probability	
AI7503000983_16205378_c1_815	NTID 425	<u>AAID</u> 4197	- gcore probability	

NO-HIT

ORF Name	NTID	AAID	NT AA LengthLength score probability
AI7503000983_162843_f2_330	426	4198	
Description		<u></u>	
sp:[LN:LYSP_ECOLI] [AC:P25737] [GN:L PERMEASE] [SP:P25737] [DB:swissprot]	YSP:CADI	R] [OR	:ESCHERICHIA COLI] [DE:LYSINE-SPECIFIC
			NT AA
ORF Name	NTID	AAID	LengthLength score probability
A17503000983_16835388_f1_19	427	4199	654 217 265 6.2e-23
Description gp:[GI:d1023735:g2547082] [LN:D89963 pho regulon] [GN:phoU] [OR:Enterobac (strain:IF03320) DNA] [DB:genpept-bc and phoU genes, complete cds.] [NT:Th [LE:4554] [RE:5279] [DI:direct]	ter cloa t1] [DE	acae] :Enter	[SR:Enterobacter cloacae obacter cloacae pstS, pstC, pstA, pstB
ORF Name	NTID	AAID	NT AA LengthLength
AI7503000983_17040911_c1_771	428	4200	123 40
Description			
NO-HIT			
ORF Name	NTID	AAID	NT AA score probability
AI7503000983_190707_±1_81	429	4201	129 42
Description			
NO-HIT			
ORF Name	NTID	AAID	NT AA score probability
AI7503000983_194431_c3_1033	430	4202	<u></u>
Description		Ĺ	J
NO-HIT			
ORF Name	NTID	AAID	NT AA score probability
AI7503000983_19540678_c1_797	431	4203	438 145 270 7.7e-23
Description		L	
<pre>gp:[GI:g4835822] [LN:AF102174] [AC:A] [GN:betL] [OR:Listeria monocytogenes] glycine betaine transporter BetL (bet [DI:direct]</pre>	DB:ge	npept	-bct2] [DE:Listeria monocytogenes
ORF Name	NTID	AAID	NT AA score probability
A17503000983_19730438_c2_926	432	4204	1272 423 2221 3.3e-230
Description			d
pir:[LN:JC5326] [AC:JC5326] [PN:meth [CL:methicillin resistance factor fer >gp:[GI:g1815620] [LN:SEU23714] [AC:U epidermidis] [DB:genpept-bct1] [DE:St methicillinresistance FEMB (femB) ger methicillin resistance] [LE:33] [RE:D	nA] [OR: J23714] caphyloc ne, comp	Staphy [PN:FI coccus clete o	ylococcus epidermidis] [DB:pir2] EMB] [GN:femB] [OR:Staphylococcus epidermidis factor essential for cds.] [NT:Factor essential for

ORF Name	NTID AAID NT AA score probability					
A17503000983_19735887_c1_754						
Description	J					
[CL:conserved hypothetical protein by sqp:[GI:e1185292:g2634073] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-by	onserved hypothetical protein ymcB] [GN:ymcB] b0835] [OR:Bacillus subtilis] [DB:pir2] 80009] [AC:Z99112:AL009126] [GN:ymcB] [FN:unknown] bct1] [DE:Bacillus subtilis complete genome (section [NT:similar to hypothetical proteins] [LE:173791]					
ORF Name	NTID AAID NT AA score probability					
A17503000983_19953281_c3_1016	434 4206 1347 448					
Description						
[DE:HOMOSERINE DEHYDROGENASE, (HDH)] [LN:BACHOM] [AC:M23217:J04034] [PN:h subtilis] [SR:Bacillus subtilis DNA]	HOM:TDM] [OR:BACILLUS SUBTILIS] [EC:1.1.1.3]] [SP:P19582] [DB:swissprot] >gp:[GI:g558494] homoserine dehydrogenase] [GN:hom] [OR:Bacillus L] [DB:genpept-bct1] [EC:1.1.1.3] [DE:Bacillus hom) gene, completecds, threonine synthase (thrC) 1577] [DI:direct]					
ORF Name	NTID AAID NT AA score probability					
A17503000983_20086512_c2_899						
Description						
	AF011784] [PN:catalase] [GN:katA] [OR:Vibrio prio fischeri catalase (katA) gene, complete cds.]					
ORF Name	NTID AAID NT AA score probability					
A17503000983_20095803_c3_994	436 4208 207 68					
Description						
NO-HIT						
ORF Name	NTID AAID NT AA score probability					
A17503000983_20312510_f3_506	437 4209 1041 346 1070 3.1e-108					
Description						
gp:[GI:g3800825] [LN:AF076684] [AC:AF076684] [PN:oligopeptide transporter putative membrane] [GN:opp-2B] [OR:Staphylococcus aureus] [DB:genpept-bct2] [DE:Staphylococcus aureus oligopeptide transporter putative membranepermease domain (opp-2B), oligopeptide transporter putativemembrane permease domain (opp-2C), oligopeptide transporterputative ATPase domain (opp-2D), and oligopeptide transporterputative ATPase domain (opp-2F) genes, complete cds.] [LE:173] [RE:1159] [DI:direct]						
ORF Name	NTID AAID NT AA score probability					
AI7503000983_20347165_f3_485	438 4210 147 48					
Description						

NO-HIT

ORF Name	NTID	AAID	<u>NT</u> LengthLe	AA score	probability
A17503000983 20370457 c2 884	439	4211	1183 6		
Description			ا لـــــالـ		
NO-HIT					
					
ORF Name	NTID	AAID	<u>NT</u> LengthLe	AA ength	probability
A17503000983_20400051_f2_319	440	4212	810 2	69 691	4.4e-68
Description sp:[LN:LEXA_BACSU] [AC:P31080] [GN:Li PROTEIN LEXA/DINR] [SP:P31080] [DB:si [PN:transcription repressor of SOS re [OR:Bacillus subtilis] [DB:pir2] >gp [OR:Bacillus subtilis] [SR:Bacillus subtilis lexA gene, 3' end.] [LE:390 [LN:BSUB0010] [AC:Z99113:AL009126] [FN:negative regulation of the SOS re [DE:Bacillus subtilis complete genome [NT:alternate gene name: dinR] [SP:Page 1.5]	wissprotegulon l :[GI:g28 subtilis] [RE:10 PN:trans egulon] e (secti	>pi exA/d 39288 DNA 007 [script [OR:B	r:[LN:A4:inR] [GN [LN:BAC] [DB:gen] DI:directional resactillus: of 21):	1315] [AC: :lexA:dinR LEXA] [AC: pept-bct1] t] >gp:[GI gulator] [subtilis] from 1781	A41315:B69651]] [CL:lexA protein] M64684] [GN:lexA] [DE:Bacillus :e1183444:g2634169] GN:lexA] [DB:genpept-bct1] 201to 2014980.]
ORF Name	NTID	AAID	<u>NT</u> LengthLe	AA ength	probability
AI7503000983_20423127_c3_991	441	4213	2676 8	91 2302	8.6e-239
pir:[LN:C69663] [AC:C69663] [PN:DNA [CL:DNA mismatch repair protein mut5] >gp:[GI:e1185295:g2634076] [LN:BSUB0] mismatch repair recognition] [OR:Bacsubtilis complete genome (section 9 [LE:176693] [RE:179269] [DI:direct]] [OR:Ba 009] [AC illus su	cillu ::Z991: ibtili:	s subtil: 12:AL009: s] [DB:ge	is] [DB:pi 126] [GN:m enpept-bct	r2] utS] [FN:DNA 1] [DE:Bacillus
ORF Name	NTID	AAID	NT LengthLe	AA score	probability
AI7503000983_20485875_c3_1008	442	4214	231 7		
Description NO-HIT			J L		
ORF Name	NTID	AAID	<u>NT</u> LengthLe	AA ength	probability
A17503000983_20501250_c2_922	443	4215	1029 34	42 551	3.0e-53
Description sp:[LN:TRPD_METJA] [AC:Q57686] [GN:TI [EC:2.4.2.18] [DE:ANTHRANILATE PHOSPI >pir:[LN:C64329] [AC:C64329] [PN:ant phosphoribosyltransferase:trpD homolo [DB:pir2] [MP:FOR225111-226121] >gp [PN:anthranilate synthase component i jannaschii] [DB:genpept-bct2] [DE:Met complete genome.] [NT:similar to GB:N [RE:4672] [DI:direct]	HORIBOSY thranila ogy] [OR :[GI:g15 II (trpD thanococ	LTRANS te syn :Metha 90969 [GI cus ja	SFERASE,] nthase, s anococcus [LN:U67 N:MJ0234] annaschii	[SP:Q5766 subunit II s jannasch: 7479] [AC:U [OR:Metha i section 2	B6] [DB:swissprot] '] [CL:anthranilate ii] [EC:4.1.3.27] J67479:L77117] anococcus 21 of 150 of the

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-bct1] 30.]
-bct1] 30.]
-bct1]

NO-HIT

ORF Name	NTID	AAID	NT AA LengthLengt)	score	probability
AI7503000983_21641877_f2_250	449	4221	954 317	687	1.2e-67
Description					
pir:[LN:E69419] [AC:E69419] [PN:pho homolog] [CL:phow protein] [OR:Archa [LN:AE001010] [AC:AE001010:AE000782] [GN:AF1358] [OR:Archaeoglobus fulgid section 97 of 172 of the complete ge GB:X02723] [LE:6184] [RE:7035] [DI:d	eoglobus [PN:pho us] [DB: nome.] [fulgi sphate genper	idus] [DB:pi e ABC transp pt-bct2] [DE	r2] >gp orter, :Archae	e:[GI:g2649217] permease protein] oglobus fulgidus
ORF Name	NTID	AAID	<u>NT AA</u> LengthLength	score	probability
A17503000983_21751938_£3_601	450	4222	708 235	183	3.0e-14
Description					
gp:[GI:g2897104] [LN:AF020798] [AC:A lipoprotein] [OR:Streptococcus therm [DE:Streptococcus thermophilus bacte putative host cell surface-exposed l Cro-like regulatoryprotein, and P1-a [NT:orf142] [LE:3941] [RE:4369] [DI:	ophilus l riophage ipoprote ntirepre	bacter lysog in,put ssor h	riophage TP- geny module, cative metal	J34] [D integr lo-prot	B:genpept-phg] asehomolog (int), einase, repressor,
ORF Name		AAID	<u>NT</u> <u>AA</u> LengthLength	score	probability
A17503000983_21912535_f1_212	451	4223	139	75	0.018
Description pir: [LN:B21124] [AC:B21124] [PN:Bkm CS314] [GN:Bkm] [OR:Drosophila mela				ion hyp	othetical protein
ORF Name	NTID	AAID	<u>NT AA</u> LengthLength	score	probability
AI7503000983_2195265_£1_6	452	4224	147 48	-]	
Description				_	
NO-HIT					
ORF Name	NTID	AAID	<u>NT AA</u> LengthLength	score	probability
A17503000983_22304635_f1_113	453	4225	150 49]	
Description					
NO-HIT	- Other Control of the Control of th	TO AN ACTION AT TABLE			
ORF Name	NTID 2	AAID	<u>NT AA</u> LengthLength	score	probability
A17503000983_22368803_c2_857	454	4226	2397 798	1659	1.2e-170
Description					
sp:[LN:SP3E_BACSU] [AC:P21458:P21459] SPORULATION PROTEIN E] [SP:P21458:P2] [AC:S09411:A32269:B32269:F69712] [P1 [OR:Bacillus subtilis] [DB:pir2] >gp [AC:Z99112:AL009126] [PN:DNA transloperatitioning through] [OR:Bacillus standard through] [OR:Bac	1459] [DE N:DNA tra :[GI:e118 case] [GE ubtilis]	B:swis ansloc B5271: N:spoI [DB:g	sprot] >pir: ase spoIIIE] g2634052] [I IIE] [FN:req enpept-bct1]	[LN:S0] [GN:s] N:BSUB: uired: [DE:B	9411] poIIIE] 0009] for chromosome acillus subtilis

[RE:155611] [DI:direct]

ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	score	probability
AI7503000983_22390917_f3_560	455	4227	165	54]	
Description						
NO-HIT				0		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
AI7503000983_22444075_c3_959	456	4228	873	290	788	2.3e-78
Description			J	<u></u>		
gp:[GI:d1034831:g3767595] [LN:AB01519] [SR:Staphylococcus aureus (strain:RN-aureus gene for LytN and Eprh, complete cds.] [SR:Staphylococcus aureus (strain:RN-gene for LytN and Eprh, complete cds.]	450) DNA ete cds. 195] [AC	A] [DB .] [NT C:AB01! A] [DB	:genpep :ORF4] 5195] [:genpep	t-bct1 [LE:32 OR:Sta ot] [DE	DE: 21] [F phyloc Staph	Staphylococcus E:4024] [DI:direct] occus aureus] ylococcus aureus
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
AI7503000983_22453425_±1_196	457	4229	138	45		
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
AI7503000983_22470463_c3_989	458	4230	810	269	881	3.3e-88
Description pir:[LN:G69884] [AC:G69884] [PN:cons [CL:hypothetical protein ymdB] [OR:Ba >gp:[GI:e1185288:g2634069] [LN:BSUB00 [OR:Bacillus subtilis] [DB:genpept-ba 9 of 21): from 1598421to 1807200.] [NE:170683] [DI:direct]	acillus 009] [AC ct1] [DE	subti ::Z991: ::Baci	lis] [D L2:AL00 Llus su	B:pir2 9126] btilis	GN:ym	dB] [FN:unknown] ete genome (section
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000983_22539812_c1_768	459	4231	1254			1.5e-131
Description						
pir:[LN:C69888] [AC:C69888] [PN:cons [CL:hypothetical protein ynbB] [OR:Ba [LN:BSU66480] [AC:U66480] [PN:YnbB] [DE:Bacillus subtilis SpoVK (spoVK), synthetase (glnA), YnaA (ynaA), YnaB (ynaF), YnaG (ynaG), YnaH(ynaH), YnaI (xynB), xylose repressor (xylR), xylos (yncB), YncC (yncC), YncD (yncD) and [RE:3839] [DI:direct] >gp:[GI:e118340] [GN:ynbB] [FN:unknown] [OR:Bacillus s	Cillus [GN:ynbE YnbA (y (ynaB), [(ynaI) se isome YncE (y 3:g2634	subtil B] [OR: ThA), YnaC; YnaC; rase (ThE)ge	is] [D Bacill YnbB ((ynaC), J (ynaJ (xylA), enes, c LN:BSU	B:pir2 us sub ynbB), YnaD), xyl xylul omplet B0010]] >gp: tilis] GlnR((ynaD) an bet ose ki e cds. [AC:Z	[GI:g1750109] [DB:genpept-bct1] glnR), glutamine , YnaE (ynaE), YnaF a-1,4-xylosidase nase(xylB), YncB] [LE:2574] 99113:AL009126]

complete genome (section 10 of 21): from 1781201to 2014980.] [NT:similar to

hypothetical proteins] [LE:94625] [RE:95890] [DI:direct]

ORF Name	NTID	AAID	NT AA LengthLength score probability	
AI7503000983_22663932_f1_22	460	4232	972 323 1021 4.8e-103	7
Description gp:[GI:g3800826] [LN:AF076684] [AC:Amembrane] [GN:opp-2C] [OR:Staphylocoaureus oligopeptide transporter putatransporter putativemembrane permeas ATPase domain (opp-2D), and oligopepgenes, complete cds.] [LE:1152] [RE:	ccus au tive men e domain tide tra	reus] mbrane n (opp anspor	[DB:genpept-bct2] [DE:Staphylococcu epermease domain (opp-2B), oligopept o-2C), oligopeptide transporterputat cterputative ATPase domain (opp-2F)	ide
ORF Name	NTID	AAID	NT AA score probability	,
A17503000983_22850885_c1_752	461	4233	1596 531 1799 1.7e-185	1
Description pir:[LN:F69884] [AC:F69884] [PN:con [CL:hypothetical protein ymdA] [OR:B >gp:[GI:e1185287:g2634068] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-b 9 of 21): from 1598421to 1807200.] [[RE:169820] [DI:direct]	acillus 009] [A0 ct1] [D1	subti C:Z991 E:Baci	llis] [DB:pir2] 12:AL009126] [GN:ymdA] [FN:unknown] Illus subtilis complete genome (sect	ion
ORF Name	NTID	AAID	NT AA score probability	
AI7503000983_23445463_c3_965	462	4234	801 266 1002 4.9e-101	1
Description				J
pir:[LN:A69699] [AC:A69699:S09561] BS1] [GN:rpsB] [CL:Escherichia coli [DB:pir2] >gp:[GI:e1185240:g2634021] protein S2] [GN:rpsB] [OR:Bacillus s complete genome (section 9 of 21): f [RE:119645] [DI:direct]	ribosom [LN:BST] ubtilis]	mal pro JB0009 [DB:	rotein S2] [OR:Bacillus subtilis] D] [AC:Z99112:AL009126] [PN:ribosoma: genpept-bct1] [DE:Bacillus subtilis	1
ORF Name	NTID	AAID	NT AA LengthLength score probability	
AI7503000983_23472175_c3_975	463	4235	417 138	1
Description sp:[LN:RBFA_BACSU] [AC:P32731] [GN:R] FACTOR A (P15B PROTEIN)] [SP:P32731] [AC:G36905:G69689:S31996] [PN:ribos] [CL:Escherichia coli protein P15B] [GI:Escherichia coli protein P15B] [GI:BSUBO] [DE:B.subtilis infB-nusA operon.] [Si:Sp:[GI:e1185256:g2634037] [LN:BSUBO] [Factor A] [GN:rbfA] [OR:Bacillus subcomplete genome (section 9 of 21): fi:ymxE, ylxO] [SP:P32731] [LE:137422]	[DB:swince] come-bince	issproding fallus sillus sillus [LE C: Z991: [DB:gen]	ct] >pir:[LN:G36905] factor A rbfA] [GN:rbfA] subtilis] [DB:pir2] >gp:[GI:g580902] subtilis] [DB:genpept-bct1] d:5131] [RE:5484] [DI:direct] d:12:AL009126] [PN:ribosome-binding enpept-bct1] [DE:Bacillus subtilis en 1807200.] [NT:alternate gene name:	8
ORF Name	NTID	AAID	NT AA score probability	
AI7503000983_23491077_c1_824	464	4236		1
Description sp:[LN:DCDA_BACSU] [AC:P23630] [GN:L' [DE:DIAMINOPIMELATE DECARBOXYLASE, (1) >gp:[GI:d1013342:g1304007] [LN:BACJHo subtilis] [SR:Bacillus subtilis (stra [DE:Bacillus subtilis DNA, 283 Kb re	YSA:LYS] DAP DECA 642] [AC ain:JH64	[OR:1 ARBOXY] C:D8443	BACILLUS SUBTILIS] [EC:4.1.1.20] [LASE)] [SP:P23630] [DB:swissprot] [32:D82370] [PN:LysA] [OR:Bacillus [C2 PheA1)) DNA] [DB:genpept-bct1]	I
[RE:282700] [DI:direct]			-	

ORF Name	NTID	AAID LengthLength score probability
A17503000983 23554760 f3 646	465	[4237][255] [84] [64][0.0077]
Description gp:[GI:g1123040] [LN:CELF44A2] [AC:U-	41993] [istol N2 o C4-typ	GN:F44A2.4] [OR:Caenorhabditis elegans]] [DB:genpept-inv1] [DE:Caenorhabditis e zinc finger and to C. elegans]
ORF Name	NTID	AAID NT AA score probability
A17503000983_23557807_f2_357 Description gp:[GI:e1172770:g2598550] [LN:LLAJ10:	466 9] [AC:A	4238 534 177 506 1.8e-48
	:genpept	-bct1] [DE:Lactococcus lactis carB and gpo
ORF Name	NTID	AAID <u>NT AA</u> LengthLength
A17503000983_23597802_c2_942	467	4239 144 47
Description NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000983_23632758_c1_723	468	4240 258 85 79 0.037
	s] [DB:g	N:phase-2 flagellin structural protein] enpept-bct2] [DE:Salmonella choleraesuis n (fljB) gene, complete cds.] [LE:1]
ORF Name	NTID	AAID NT AA score probability
AI7503000983_23633467_c2_851	469	4241 987 328 659 1.1e-64
[CL:conserved hypothetical protein H3 >gp:[GI:e269877:g1592690] [LN:BSRIBRE [GN:ribC] [OR:Bacillus subtilis] [DB: genes.] [NT:riboflavin kinase] [SP:P5 >gp:[GI:e1185258:g2634039] [LN:BSUB00] [GN:ribC] [FN:riboflavin biosynthesis	[0963] [PS] [AC: genpept 54575] [009] [AC] [OR:B	Z80835] [PN:FMN adenylyltransferase] -bct1] [DE:B.subtilis ribC, rpsO and pnpA LE:88] [RE:1038] [DI:direct] :Z99112:AL009126] [PN:FAD synthase]

[DI:direct]

NT ORF Name score probability Length Length A17503000983 23650250 c1 729 181 1.6e-65 4242 546 Description sp:[LN:HSLV BACSU] [AC:P39070] [GN:HSLV:CLPQ:CODW] [OR:BACILLUS SUBTILIS] [EC:3.4.99.-] [DE:HEAT SHOCK PROTEIN HSLV PRECURSOR,] [SP:P39070] [DB:swissprot] >pir:[LN:S61494] [AC:S61494:S45024:C69601] [PN:20S proteasome beta-type chain clpQ:heat shock protein codW] [GN:clpQ:codW] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:g535349] [LN:BSU13634] [AC:U13634] [PN:CodW] [GN:codW] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis JH642 dipeptide permease operon regulators, codV,codW, codX, and codY genes, complete cds.] [LE:1220] [RE:1765] [DI:direct] >gp:[GI:e1185206:g2633987] [LN:BSUB0009] [AC:Z99112:AL009126] [PN:beta-type subunit of the 20S proteasome] [GN:clp0] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 9 of 21): from 1598421to 1807200.] [NT:alternate gene name: hslV, codW] [SP:P39070] [LE:89093] [RE:89638] [DI:direct] NT AΑ ORF Name NTID AAID <u>score</u> probability LengthLength A17503000983 23650343 c2 949 1167 388 1.1e-50 Description pir:[LN:G71097] [AC:G71097] [PN:probable amidohydrolase] [GN:PH1043] [OR:Pyrococcus horikoshii] [DB:pir2] >qp:[GI:d1031084:q3257458] [LN:AP000004] [AC:AP000004:AB009494:AB009495:AB009496:AB009497:AB009498:AB009499] [PN:387aa long hypothetical amidohydrolase] [GN:PH1043] [OR:Pyrococcus horikoshii] [SR:Pyrococcus horikoshii (strain:OT3) DNA] [DB:genpept-bct1] [DE:Pyrococcus horikoshii OT3 genomic DNA, 777001-994000 nt. position(4/7).] [NT:similar to Swiss Prot:P80092 percent identity:] [LE:172136] [RE:173299] [DI:complement] AΑ ORF Name NTID AAID probability score LengthLength A17503000983 23671890 cl 807 472 1287 428 3.9e-92 918 Description pir:[LN:A69730] [AC:A69730] [PN:UV-damage repair protein uvrX] [GN:uvrX] [CL:umuC protein] [OR:Bacillus subtilis] [DB:pir2] >qp:[GI:e1183597:q2634570] [LN:BSUB0012] [AC:Z99115:AL009126] [PN:UV-damage repair protein] [GN:uvrX] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 12 of 21): from 2195541to 2409220.] [NT:alternate gene name: yolE] [LE:74108] [RE:75358] [DI:complement] >gp:[GI:g3025495] [LN:AF020713] [AC:AF020713] [PN:IMPB/MUCB/SAMB family protein] [GN:yolE] [OR:Bacteriophage SPBc2] [DB:genpept-phg] [DE:Bacteriophage SPBc2 complete genome.] [LE:14792] [RE:16042] [DI:direct] NTAA ORF Name NTID <u>score</u> AAID probability LengthLength A17503000983_23673150_c3_1029 4245 473 1002 1.1e-135 Description pir:[LN:C70015] [AC:C70015] [PN:GMP reductase, homolog yumD] [GN:yumD] [OR:Bacillus

subtilis] [EC:1.6.6.8] [DB:pir2] >gp:[GI:e1184292:g2635710] [LN:BSUB0017] [AC:Z99120:AL009126] [GN:yumD] [FN:unknown] [OR:Bacillus subtilis] [DB:qenpept-bct1] [DE:Bacillus subtilis complete genome (section 17 of 21): from 3197001to 3414420.] [NT:similar to GMP reductase] [LE:105099] [RE:106079] [DI:direct] >gp:[GI:e311468:g1934831] [LN:BSZ93939] [AC:Z93939] [PN:unknown] [GN:yumD] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:B.subtilis genomic DNA fragment from yumA to yulF.] [NT:potential inosine or guanosine 5' monophosphate] [LE:3536] [RE:4516] [DI:direct]

RF Name NTID AAID LengthLength score probability
17503000983 23725442 £2 230 474 4246 456 151 197 10.0028
escription
p:[GI:g3582212] [LN:AE001272] [AC:AE001272] [PN:Potential membrane spanning protein] GN:ORF00031] [OR:Lactococcus lactis] [DB:genpept-bct2] [DE:Lactococcus lactis DPC3147 lasmid pMRC01, complete plasmidsequence.] [NT:similar to GB:Z30588 PID:459257 percent dentity:] [LE:25025] [RE:25804] [DI:complement]
RF Name NTID AAID NT AA score probability
17503000983_23860952_f3_526
p:[LN:PMSR_BACSU] [AC:P54154] [GN:YPPP] [OR:BACILLUS SUBTILIS] [DE:REDUCTASE)] SP:P54154] [DB:swissprot] >pir:[LN:E69940] [AC:E69940] [PN:peptide methionine ulfoxide reductase homolog yppP] [GN:yppP] [CL:peptide methionine sulfoxide eductase] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:g1256653] [LN:BACYACA] [AC:L77246] PN:DNA-binding protein] [GN:yppP] [OR:Bacillus subtilis] [DB:genpept-bct1] DE:Bacillus subtilis (YAC10-9 clone) DNA region between the serA andkdg loci.] NT:42.4% identity with the Lycopersicon esculentum] [LE:33389] [RE:33922] [DI:direct] gp:[GI:e1183616:g2634589] [LN:BSUB0012] [AC:Z99115:AL009126] [GN:yppP] [FN:unknown] OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 2 of 21): from 2195541to 2409220.] [NT:similar to peptide methionine sulfoxide eductase] [SP:P54154] [LE:91230] [RE:91763] [DI:complement]
NTID AAID NT AA Score probability 17503000983_23944052_c1_789 476 4248 378 125 248 1.7e-20
escription
p:[LN:CATB_BACFR] [AC:P45737] [GN:KATB] [OR:BACTEROIDES FRAGILIS] [EC:1.11.1.6] DE:CATALASE,] [SP:P45737] [DB:swissprot] >pir:[LN:A57262] [AC:A57262] [PN:catalase,] GN:katB] [CL:catalase] [OR:Bacteroides fragilis] [EC:1.11.1.6] [DB:pir2] gp:[GI:g841192] [LN:BFU18676] [AC:U18676] [PN:catalase] [GN:katB] [FN:decomposes ydrogen peroxide in water and] [OR:Bacteroides fragilis] [DB:genpept-bct2] EC:1.11.1.6] [DE:Bacteroides fragilis catalase (katB) gene, complete cds.] [LE:348] RE:1808] [DI:direct]
DE:CATALASE,] [SP:P45737] [DB:swissprot] >pir:[LN:A57262] [AC:A57262] [PN:catalase,] GN:katB] [CL:catalase] [OR:Bacteroides fragilis] [EC:1.11.1.6] [DB:pir2] gp:[GI:g841192] [LN:BFU18676] [AC:U18676] [PN:catalase] [GN:katB] [FN:decomposes ydrogen peroxide in water and] [OR:Bacteroides fragilis] [DB:genpept-bct2] EC:1.11.1.6] [DE:Bacteroides fragilis catalase (katB) gene, complete cds.] [LE:348]
DE:CATALASE,] [SP:P45737] [DB:swissprot] >pir:[LN:A57262] [AC:A57262] [PN:catalase,] GN:katB] [CL:catalase] [OR:Bacteroides fragilis] [EC:1.11.1.6] [DB:pir2] gp:[GI:g841192] [LN:BFU18676] [AC:U18676] [PN:catalase] [GN:katB] [FN:decomposes ydrogen peroxide in water and] [OR:Bacteroides fragilis] [DB:genpept-bct2] EC:1.11.1.6] [DE:Bacteroides fragilis catalase (katB) gene, complete cds.] [LE:348] RE:1808] [DI:direct]
DE:CATALASE,] [SP:P45737] [DB:swissprot] >pir:[LN:A57262] [AC:A57262] [PN:catalase,] GN:katB] [CL:catalase] [OR:Bacteroides fragilis] [EC:1.11.1.6] [DB:pir2] gp:[GI:g841192] [LN:BFU18676] [AC:U18676] [PN:catalase] [GN:katB] [FN:decomposes ydrogen peroxide in water and] [OR:Bacteroides fragilis] [DB:genpept-bct2] EC:1.11.1.6] [DE:Bacteroides fragilis catalase (katB) gene, complete cds.] [LE:348] RE:1808] [DI:direct] RF Name NTID AAID NT AA LengthLength score probability
DE:CATALASE,] [SP:P45737] [DB:swissprot] >pir:[LN:A57262] [AC:A57262] [PN:catalase,] GN:katB] [CL:catalase] [OR:Bacteroides fragilis] [EC:1.11.1.6] [DB:pir2] gp:[GI:g841192] [LN:BFU18676] [AC:U18676] [PN:catalase] [GN:katB] [FN:decomposes ydrogen peroxide in water and] [OR:Bacteroides fragilis] [DB:genpept-bct2] EC:1.11.1.6] [DE:Bacteroides fragilis catalase (katB) gene, complete cds.] [LE:348] RE:1808] [DI:direct] RF Name NTID AAID NT AA LengthLength LengthLength LengthLength Escription
DE:CATALASE,] [SP:P45737] [DB:swissprot] >pir:[LN:A57262] [AC:A57262] [PN:catalase,] GN:katB] [CL:catalase] [OR:Bacteroides fragilis] [EC:1.11.1.6] [DB:pir2] gp:[GI:g841192] [LN:BFU18676] [AC:U18676] [PN:catalase] [GN:katB] [FN:decomposes ydrogen peroxide in water and] [OR:Bacteroides fragilis] [DB:genpept-bct2] EC:1.11.1.6] [DE:Bacteroides fragilis catalase (katB) gene, complete cds.] [LE:348] RE:1808] [DI:direct] RF Name NTID AAID LengthLength 17503000983_23992812_c1_793 AATO O-HIT RF Name NTID AAID NT AA Score probability
DE:CATALASE, [SP:P45737] [DB:swissprot] >pir:[LN:A57262] [AC:A57262] [PN:catalase,] GN:katB [CL:catalase] [OR:Bacteroides fragilis] [EC:1.11.1.6] [DB:pir2] gp:[GI:g841192] [LN:BFU18676] [AC:U18676] [PN:catalase] [GN:katB] [FN:decomposes ydrogen peroxide in water and] [OR:Bacteroides fragilis] [DB:genpept-bct2] EC:1.11.1.6] [DE:Bacteroides fragilis catalase (katB) gene, complete cds.] [LE:348] RE:1808] [DI:direct] RF Name NTID AAID NT AA Score probability NTID AAID NT AA AA

OKT Name	MIID	AAID	Length	Length	BCOLE	probability
A17503000983_24033217_c2_850	479	4251	930	309	720	3.8e-71
Description sp:[LN:TRUB_BACSU] [AC:P32732] [GN:T. [DE:HYDROLYASE)] [SP:P32732] [DB:swing protein P35] [OR:Bacillus subtilis] [AC:Z99112:AL009126] [PN:tRNA pseudorsubtilis] [DB:genpept-bct1] [DE:Bacilfrom 1598421to 1807200.] [NT:alternative:138786] [DI:direct]	ssprot] ruB:P35 [DB:pir2 uridine llus sul	>pir: homolo 2] >gp: 55 syr otilis	[LN:G69 og] [GN :[GI:e1 othase] comple	9726] [J:truB .185257 [GN:t ete gen	AC:G69] [CL: :g2634 ruB] ome (s	P726:H36905:S31997] Escherichia coli P038] [LN:BSUB0009] P0R:Bacillus Rection 9 of 21):
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
AI7503000983_24078753_c3_999	480	4252	450	149	295	4.1e-26
Description pir: [LN:G69657] [AC:G69657] [PN:tRN [GN:miaA] [CL:delta(2)-isopentenylp [DB:pir2] >gp:[GI:e1183392:g2634117] isopentenylpyrophosphate transferase [DB:genpept-bct1] [DE:Bacillus subtil 1781201to 2014980.] [LE:84430] [RE:89	yrophosm [LN:BSU] [GN:mi lis comm	phate t JB0010] iaA] [C plete g	ransfe [AC:Z R:Baci genome	rase] 399113: llus s	[OR:Ba AL0091 ubtili	cillus subtilis] 26] [PN:tRNA s]
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
AI7503000983_24222137_£2_341	481	4253	159	52		
Description			·			
NO-HIT						
ORF Name AI7503000983_24225000_f3_475	NTID 482	<u>AAID</u> 4254	<u>NT</u> Length	AA Length 297	score	probability 5.3e-33
Description	L		l L	L	L	
gp:[GI:g2194195] [LN:SGU61158] [AC:U6 gallinarum] [DB:genpept-bct1] [DE:Sta putative membraneprotein (gdmH), ABC galliderminprecursor (gdmA) genes, co modifying enzyme (gdmB) genes, partia (ATP-binding] [LE:179] [RE:874] [DI:	aphyloco transpo omplete al cds.]	occus g orter (cds, p [NT:p	allina gdmT), outativ	rum Tu and a e memb	e3928 ntibio rane p	GdmF (gdmF), tic rotein(gdmE) and
ORF Name	NTID	<u>AAID</u>	<u>NT</u> Length	<u>AA</u> Length	score	probability
AI7503000983_24225053_c3_1031	483	4255	318	105	202	2.9e-16
Description pir:[LN:E69894] [AC:E69894] [PN:hypound	g263417 Subtilis	72] [LN 3] [DB:	:BSUB0 genpep	010] [2 t-bct1]	AC:Z99] [DE:	113:AL009126] Bacillus subtilis
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
AI7503000983_24251466_c1_799	484	4256	2745		3437	0.0
Description		L	·			
sp:[LN:ACON_BACSU] [AC:P09339:Q45059] [DE:ACONITATE HYDRATASE, (CITRATE HYI [DB:swissprot]						

ORF Name A17503000983_24257252_c1_727	NTID	AAID NT AA score probability [4257 177 58]				
Description NO-HIT						
ORF Name	NTID	AAID NT AA score probability				
A17503000983_24267941_c1_745	486	4258 726 241 304 4.5e-27				
[DB:swissprot] >pir:[LN:B69885] [AC:homolog ymfC] [GN:ymfC] [OR:Bacillu [LN:BSUB0009] [AC:Z99112:AL009126] [DB:genpept-bct1] [DE:Bacillus subti	B69885 s subti GN:ymfC lis com	R:BACILLUS SUBTILIS] [DE:REGION] [SP:O31761]] [PN:transcription regulator GntR family lis] [DB:pir2] >gp:[GI:e1185272:g2634053]] [FN:unknown] [OR:Bacillus subtilis] plete genome (section 9 of 21): from ptional regulator (GntR family)] [SP:O31761]				
ORF Name	NTID	AAID NT AA probability LengthLength score probability				
A17503000983_24267942_c1_823	487	4259 1089 362 300 1.2e-26				
[DE:ALANINE RACEMASE,] [SP:P10725:P9 [LN:AB001488] [AC:AB001488] [PN:ALAN [SR:Bacillus subtilis (strain:168) D subtilis genome sequence, 148 kb seq [LE:50435] [RE:51604] [DI:direct] >g [AC:Z99106:AL009126] [PN:D-alanine r	6620] [1 INE RACI NA] [DB uence o: p:[GI:e: acemase acillus	[GN:dal] [OR:Bacillus subtilis] subtilis complete genome (section 3 of 21):				
ORF Name	NTID	AAID <u>NT AA</u> score probability				
AI7503000983_24275017_c2_856	488	4260 144 47				
Description NO-HIT						
ORF Name	NTID	AAID NT AA score probability				
A17503000983_24352200_f3_600	489	4261 1095 364 674 2.8e-66				
Description pir:[LN:T02833] [AC:T02833] [PN:hypothetical protein L4171.5] [OR:Leishmania major] [DB:pir2] [MP:1] >gp:[GI:g2266911] [LN:AE001274] [AC:AE001274:AC003011:AC002552:U60409:AF008205:AC002134:AF008206:U7 0253] [PN:L4171.5] [GN:L4171.5] [OR:Leishmania major] [DB:genpept-inv2] [DE:Leishmania major chromosome 1, complete sequence.] [NT:similar to threonine aldolase; leucine zipper] [LE:124709] [RE:125788] [DI:direct]						
ORF Name	NTID	AAID NT AA score probability				
A17503000983_24407758_t3_696	490	4262 129 42				
Description NO-HIT						

ORF Name	NTID	AAID LengthLength score probability
AI7503000983_24412811_c3_997	491	4263 957 318 147 4.4e-11
<pre>sativa] [SR:, rice] [DB:pir2] >gp:[G</pre>	I:g2801 PL1] [O	DR:Oryza sativa] [DB:genpept-pln2] [DE:Oryza
ORF Name	NTID	AAID NT AA score probability
AI7503000983_24414187_c3_1003	492	4264 396 131 74 0.011
Description gp:[GI:g42727] [LN:ECRHAT] [AC:X6069 [DB:genpept-bct1] [DE:E. coli rhaT g sodA(5' end) genes.] [SP:P00448] [LE	ene for	L-rhamnose permease, rhaC (3' end) and
ORF Name	NTID	AAID <u>NT AA</u> LengthLength score probability
A17503000983_24414818_c1_738	493	4265 870 289 952 9.8e-96
	acillus	C:Z99112:AL009126] [GN:nusA] s subtilis] [DB:genpept-bct1] [DE:Bacillus from 1598421to 1807200.] [LE:133252]
ORF Name	NTID	AAID LengthLength score probability
A17503000983_24415933_c3_1078 Description NO-HIT	494	4266 153 51
ORF Name A17503000983_24417512_f3_505 Description NO-HIT	<u>NTID</u> 495	AAID NT AA score probability 4267 135 44
ORF Name A17503000983_24470927_f2_358 Description NO-HIT	NTID 496	AAID NT AA score probability 4268 138 45
ORF Name A17503000983_24475252_c3_1052 Description NO-HIT	NTID 497	AAID NT AA score probability 4269 126 41
ORF Name A17503000983_24484683_c3_1041 Description	NTID 498	AAID NT AA score probability 4270 129 42
NO-HIT		

ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	score	probability
AI7503000983_24492827_f1_223	499	4271	2601	866	1069	3.9e-108
Description			JLJ	<u> </u>	J L	
pir:[LN:G69801] [AC:G69801] [PN:hypsubtilis] [DB:pir2] >gp:[GI:e1182850 [GN:yfhO] [FN:unknown] [OR:Bacillus complete genome (section 5 of 21): f[DI:direct] >gp:[GI:d1025397:g280454 subtilis] [SR:Bacillus subtilis DNA] genome sequence, 79 to 81 degree reg	:g263318 subtilis rom 8028 5] [LN:D [DB:gen	4] [L] [DB 21 to 85082 pept-	N:BSUBO :genper 1011250] [AC:I bct1] [005] [bt-bct1].] [LE 085082] [DE:Bac	AC: Z9: DE: 1286: PN: S: illus	9108:AL009126] :Bacillus subtilis 91] [RE:131150] /fhO] [OR:Bacillus subtilis DNA,
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
AI7503000983_24495928_c1_772 Description	500	4272	210	69	70	0.028
pir:[LN:G69058] [AC:G69058] [PN:hyp [OR:Methanobacterium thermoautotroph [AC:AE000905:AE000666] [PN:unknown] thermoautotrophicum] [DB:genpept-bct bases 1293944 to 1305586(section 111 Code:14.00 - Unknown, ; similar to,]	icum] [D [GN:MTH1 2] [DE:M of 148)	B:pir 440] ethan of t	2] >gp: [OR:Met obacter he comp	[GI:g2 hanoba ium th lete g	622552 cteriu ermoau	2] [LN:AE000905] um utotrophicum from .] [NT:Function
ORF Name	NTID	<u>AAID</u>	<u>NT</u> Length	<u>AA</u> Length	score	probability
AI7503000983_2459667_c1_796	501	4273	174	57		
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	score	probability
A17503000983_24631637_c3_1002	502	4274	708	235	236	7.3e-20
Description			<u> </u>		<u></u>	
gp:[GI:g2444107] [LN:U88974] [AC:U88 temperate bacteriophage O1205] [DB:g temperate bacteriophage O1205, compl	enpept-p	hg] [1	DE:Stre	ptococ	cus th	nermophilus
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	<u>score</u>	probability
AI7503000983_24640925_c3_1037	503	4275	1272	423	1288	2.4e-131
Description						
<pre>sp:[LN:OPUD_BACSU] [AC:P54417] [GN:O TRANSPORTER OPUD] [SP:P54417] [DB:sw betaine transporter opuD] [GN:opuD]</pre>		>pir			[AC:G6	

ORF Name	NTID	AAID	NT	<u>AA</u>	score	probability
			Length	Length		
A17503000983_24647176_c2_928	504	4276	771	256	128	4.7e-06
Description gp:[GI:g4981173] [LN:AE001738] [AC:R protein] [GN:TM0651] [OR:Thermotoga section 50 of 136 of the complete ge 52.43;] [LE:13445] [RE:14251] [DI:co	maritin	na] [DB (NT:si	:genpep	t-bct2] [DE:	Thermotoga maritima
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000983_24647182_c1_821	505	4277	729	242	232	3.9e-33
Description gp:[GI:g4982086] [LN:AE001799] [AC:Areductase] [GN:TM1520] [OR:Thermotogmaritima section 111 of 136 of the cidentity: 62.67;] [LE:16971] [RE:176	ga marit complete	ima] [genom	DB:genp e.] [NT	ept-bc	t2] [I	E:Thermotoga
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000983_24659382_c2_890	506	4278	126	41		
Description	<u> </u>	J L	<u></u>			
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000983_24666043_c1_803	507	4279	1233	410	866	1.3e-86
Description			L		Ш.	
pir:[LN:D70006] [AC:D70006] [PN:conserved hypothetical protein yubA] [GN:yubA] [CL:Bacillus subtilis conserved hypothetical protein yueF] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1185989:g2635600] [LN:BSUB0016] [AC:Z99119:AL009126] [GN:yubA] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 16 of 21): from 2997771to 3213410.] [NT:similar to hypothetical proteins] [LE:196858] [RE:198024] [DI:complement]						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000983_24744010_c3_1028 Description	508	4280	273	90	347	1.3e-31
pir:[LN:F69835] [AC:F69835] [PN:rik [CL:Escherichia coli ribosomal prote >gp:[GI:e1182877:g2633211] [LN:BSUBC [OR:Bacillus subtilis] [DB:genpept-k 5 of 21): from 802821 to1011250.] [N	in S14] 005] [A oct1] [D	OR:Baci. C:Baci.	acillus 08:AL00 llus su	subti 9126] btilis	lis] [[GN:yh compl	DB:pir2] zA] [FN:unknown] ete genome (section

[RE:162867] [DI:direct]

NT AAORF Name AAID NTID score probability Length Length A17503000983 24792776 c2 846 509 528 3.6e-34 Description sp:[LN:YLXS BACSU] [AC:P32726] [GN:YLXS] [OR:BACILLUS SUBTILIS] [DE:HYPOTHETICAL 17.6 KD PROTEIN IN NUSA 5'REGION (P15A) (ORF1)] [SP:P32726] [DB:swissprot] >pir:[LN:B36905] [AC:B36905:E69882:S31990] [PN:conserved hypothetical protein ylxS] [GN:ylxS] [CL:nus operon 15K protein] [OR:Bacillus subtilis] [DB:pir2] >qp:[GI:q49315] [LN:BSORF1T7A] [AC:Z18631] [GN:ORF1] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:B.subtilis infB-nusA operon.] [SP:P32726] [LE:456] [RE:926] [DI:direct] >gp:[GI:e1185250:g2634031] [LN:BSUB0009] [AC:Z99112:AL009126] [GN:ylxS] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 9 of 21): from 1598421to 1807200.] [NT:alternate gene name: ymxA; similar to hypothetical] [SP:P32726] [LE:132747] [RE:133217] [DI:direct] NT AA ORF Name NTID AAID score probability LengthLength A17503000983 24823377 f3 498 510 912 303 8.2e-44 Description pir:[LN:E69840] [AC:E69840] [PN:hypothetical protein yitL] [GN:yitL] [OR:Bacillus subtilis] [DB:pir2] >qp:[GI:e1183105:q2633439] [LN:BSUB0006] [AC:Z99109:AL009126] [GN:yitL] [FN:unknown] [OR:Bacillus subtilis] [DB:qenpept-bct1] [DE:Bacillus subtilis complete genome (section 6 of 21): from 999501 to1209940.] [LE:181473] [RE:182369] [DI:direct] NT AΑ ORF Name NTID AAID score probability LengthLength AI7503000983_24877312_f1_105 511 4283 1392 463 1093 1.1e-110 Description pir:[LN:A69763] [AC:A69763] [PN:homoserine dehydrogenase homolog yclM] [GN:yclM] [CL:aspartate kinase homology] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1182346:g2632680] [LN:BSUB0003] [AC:Z99106:AL009126] [GN:yclM] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 3 of 21): from 402751 to611850.] [NT:similar to homoserine dehydrogenase] [LE:27453] [RE:28817] [DI:complement] >gp:[GI:d1009646:q1805449] [LN:D50453] [AC:D50453] [PN:homologue of aspartokinase 2 alpha and beta] [GN:yclM] [OR:Bacillus subtilis] [SR:Bacillus subtilis (strain:168 trpC2) DNA] [DB:genpept-bct1] [DE:Bacillus subtilis DNA for 25-36 degree region containing theamyE-srfA region, complete cds.] [LE:109856] [RE:111220] [DI:complement] NT AA ORF Name NTID score AAID probability LengthLength AI7503000983_24886550_c1_741 512 4284 288 6.1e-32 Description pir:[LN:F69700] [AC:F69700:S11365:S70690] [PN:ribosomal protein S15 (rps0):ribosomal protein BS18] [GN:rps0] [CL:Escherichia coli ribosomal protein S15: eubacterial ribosomal protein S15 homology] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e269878:g1592691] [LN:BSRIBRPS] [AC:Z80835] [PN:ribosomal protein S15] [GN:rpsO] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:B.subtilis ribC, rpsO and pnpA

genes.] [SP:P21473] [LE:1195] [RE:1464] [DI:direct] >gp:[GI:e1185259:g2634040] [LN:BSUB0009] [AC:Z99112:AL009126] [PN:ribosomal protein S15 (BS18)] [GN:rps0] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 9 of 21): from 1598421to 1807200.] [SP:P21473] [LE:139912] [RE:140181] [DI:direct]

ORF Name	NTID	<u>AAID</u>	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000983_24886677_c3_971	513	4285	1722	573		1.3e-194
Description			لــــالـ		لــــال	
pir:[LN:G69682] [AC:G69682] [PN:pro [GN:proS] [CL:prolinetRNA ligase] >gp:[GI:e1185248:g2634029] [LN:BSUBO synthetase] [GN:proS] [OR:Bacillus s complete genome (section 9 of 21): f [DI:direct]	OR:Ba O009] [A Subtilis	cillus C:Z991] [DB:	subtil 12:AL00 genpept	lis] [E 09126] :-bct1]	:C:6.1 PN:pr DE:E	[DB:pir2] colyl-tRNA Bacillus subtilis
ORF_Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000983_24900332_c2_870	514	4286	981	326	935	6.2e-94
Description		<u> </u>				
repair mutL] [GN:mutL] [CL:mismatch [DB:pir2] >gp:[GI:g1002521] [LN:BSU2 [FN:mismatch-repair recognition] [OR subtilis spore coat protein (cotE) g proteins (mutS) and (mutL) genes,com >gp:[GI:e1185296:g2634077] [LN:BSUB0 mismatch repair] [OR:Bacillus subtil complete genome (section 9 of 21): f [RE:181168] [DI:direct]	:7343] [::Bacill gene, pa mplete c [009] [A .is] [DB	AC:U27 us sub rtial ds.] [C:Z991 :genpe	343] [E tilis] cds, ar LE:2782 12:AL00 pt-bct1	PN:MutI [DB:ge idmisma] [RE:]9126] .] [DE:	[GN: npept- tch re 4665] [GN:mu Bacill	mutL] bctl] [DE:Bacillus pair recognition [DI:direct] tL] [FN:DNA us subtilis
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
AI7503000983_25445253_c3_1039	515	4287	495	164	336	1.8e-30
Description				L		
pir:[LN:F69891] [AC:F69891] [PN:con [OR:Bacillus subtilis] [DB:pir2] >gp [PN:YneP] [GN:yneP] [OR:Bacillus subtilis] the fragment; 170 degree region).] [L >gp:[GI:e1183462:g2634187] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-b10 of 21): from 1781201to 2014980.] [RE:149254] [DI:direct]	o:[GI:e2 btilis] bE:12510 0010] [A oct1] [D	49652: [DB:ge:] [RE: C:Z991 E:Baci	g140545 npept-b 12875] 13:AL00 llus su	66] [LN oct1] [[DI:di 9126] btilis	EBC170 DE:B.s rect] [GN:yn compl	DEGR] [AC:Z73234] Subtilis DNA (26.2 DEP] [FN:unknown] ete genome (section
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000983_25449061_c1_780	516	4288	249	82	191	4.3e-15
Description pir: [LN:D69901] [AC:D69901] [PN:two [GN:yocG] [CL:regulatory protein co subtilis] [DB:pir2] >gp:[GI:g2619014 [GN:yocG] [OR:Bacillus subtilis] [DB region between terC and odhAB.] [NT: [LE:71699] [RE:72298] [DI:direct] >g [AC:Z99114:AL009126] [GN:yocG] [FN:u	mA:respo] [LN:Al :genpept similar p:[GI:e:	onse re F027868 t-bct1; to B.8	egulato 3] [AC: [DE:B subtili 2:g2634	r homo AF0278 acillu s Spo0 313] [logy] 68] [P s subt A prot LN:BSU	[OR:Bacillus N:sensor regulator] ilis chromosome ein (267 aa)] B0011]

192

[DE:Bacillus subtilis complete genome (section 11 of 21): from 2000171to 2207900.] [NT:similar to two-component response regulator [YocF]] [LE:90790] [RE:91389]

[DI:direct]

ORF Name A17503000983_25578140_f3_612	<u>NTID</u>	AAID 4289	NT AA LengthLength score probability 132 43			
Description NO-HIT						
ORF Name	NTID	AAID	NT AA LengthLength score probability			
A17503000983_25587942_c3_956	518	4290	786 261 556 9.0e-54			
<pre>Description pir:[LN:C69693] [AC:C69693] [PN:rib [OR:Bacillus subtilis] [DB:pir2] >gp [AC:Z99112:AL009126] [PN:ribonucleas [DB:genpept-bct1] [EC:3.1.26.4] [DE: 21): from 1598421to 1807200.] [LE:78</pre>	:[GI:e1 e H] [G Bacillu	185197 N:rnh] s subt	:g2633978] [LN:BSUB0009] [OR:Bacillus subtilis] ilis complete genome (section 9 of			
ORF Name	NTID	AAID	NT AA score probability			
A17503000983_25593925_f1_166	519	4291	153 50			
Description NO-HIT						
ORF Name	NTID	AAID	NT AA LengthLength score probability			
A17503000983_25626625_c2_891	520	4292	636 211			
Description NO-HIT						
ORF Name	NTID	AAID	NT AA LengthLength score probability			
A17503000983_25664512_c2_915	521	4293	495 164 279 2.0e-24			
Description pir:[LN:A69805] [AC:A69805] [PN:hypothetical protein yfiW] [GN:yfiW] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1182832:g2633166] [LN:BSUB0005] [AC:Z99108:AL009126] [GN:yfiW] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 5 of 21): from 802821 to1011250.] [LE:113464] [RE:114240] [DI:direct] >gp:[GI:d1025379:g2804527] [LN:D85082] [AC:D85082] [PN:YfiW] [OR:Bacillus subtilis] [SR:Bacillus subtilis DNA] [DB:genpept-bct1] [DE:Bacillus subtilis DNA, genome sequence, 79 to 81 degree region.] [LE:6355] [RE:7131] [DI:direct]						
ORF Name	NTID	AAID	NT AA LengthLength score probability			
A17503000983_25665937_c3_968	522	4294	558 185 841 5.7e-84			
Description sp:[LN:RRF_STAAU] [AC:O33276] [GN:FR RECYCLING FACTOR (RIBOSOME RELEASING >gp:[GI:g2645713] [LN:AF033018] [AC: [OR:Staphylococcus aureus] [DB:genperecycling factor (frr) gene,complete	FACTOR) AF033018 pt-bct2]	(RRF) 3] [PN DE:	ribosome recycling factor] [GN:frr] Staphylococcus aureus ribosome			

ORF Name	NTID	AAID	NT Length	AA Length	score	probability
A17503000983_257837_c3_1013	523	4295	174	57	87	0.0045
Description gp:[GI:g2668605] [LN:AF015453] [AC:R [DB:genpept-bct2] [DE:Lactobacillus gene,partial cds; GNTR transcription genes, complete cds.] [NT:3.0E-ORF-1	rhamnos nal regu	us 6-pl lator l	hospho homolog	-beta-gi	lucosi urface	dase homolog
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
AI7503000983_25977318_c3_1044	524	4296	2415	804	3470	0.0
sp:[LN:PARC_STAAU] [AC:P50073:P95682 [EC:5.99.1] [DE:TOPOISOMERASE IV S >gp:[GI:d1011747:g1777321] [LN:D6707 subunit] [GN:grlA] [OR:Staphylococcu (isolate:RN4220) DNA] [DB:genpept-botopoisomerase IV GrlB subunit,DNA to [LE:2376] [RE:4778] [DI:direct]	SUBUNIT [AC: s aureu t1] [DE	A,] [S] D67075] s] [SR :Staphy	P:P5007 [PN:I :Staphy ylococo	73:P9568 DNA topo vlococci cus aure	32:P95 Disome us aur Eus DN	[683] [DB:swissprot] erase IV GrlA reus NA for DNA
ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	score	probability
AI7503000983_26205387_f2_331	525	4297	336	111		
<u>Description</u>						
NO-HIT	W. W. J. S. S	**************************************		100		
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000983_26210925_c3_1032	526	4298	2028	675	2097	4.5e-217
Description sp:[LN:TKT_BACSU] [AC:P45694] [GN:TK [DE:TRANSKETOLASE,] [SP:P45694] [DB: [PN:transketolase, tkt] [GN:tkt] [Cdomain homology] [OR:Bacillus subtil >gp:[GI:e249642:g1405446] [LN:BC170D [OR:Bacillus subtilis] [DB:genpept-bdegree region).] [SP:P45694] [LE:153 [LN:BSUB0010] [AC:Z99113:AL009126] [OR:Bacillus subtilis] [DB:genpept-bgenome (section 10 of 21): from 1781 [SP:P45694] [LE:137916] [RE:139919]	swisspronum (Superscript State of Superscript Superscr	ot] >pi ketolas :2.2.1. C:Z7323 E:B.suk 3540] sketola C:2.2.1	ir:[LN: se:thia 1] [DE 34] [PN otilis [DI:dir ase] [G	G69723] mine py s:pir2] I:transk DNA (26 Tect] >c N:tkt] DE:Bacil	[AC: vropho setola 5.2 kb gp:[GI [FN:p	G69723:S57401] sphate-binding se] [GN:tktA] fragment; 170 :e1183448:g2634173] entose phosphate] ubtilis complete
ORF Name AI7503000983 26213890_c3 1026	NTID	<u>AAID</u>	<u>NT</u> Length	Lengtn-	core	probability
Description	52,	1277	للتكال	232	757	I.JG-1J
gp:[GI:e313391:g2052219] [LN:SCDNACA [OR:Streptomyces coelicolor] [DB:gen [RE:1858] [DI:direct]						

 $rac{ ext{ORF Name}}{ ext{ORF Name}}$ $rac{ ext{NTID}}{ ext{LengthLength}} rac{ ext{Score}}{ ext{probability}}$

A17503000983_26257806_c3_993

528

984

327

Description

sp:[LN:MUTL_BACSU] [AC:P49850] [GN:MUTL] [OR:BACILLUS SUBTILIS] [DE:DNA MISMATCH REPAIR PROTEIN MUTL] [SP:P49850] [DB:swissprot] >pir:[LN:A69663] [AC:A69663] [PN:DNA mismatch repair mutL] [GN:mutL] [CL:mismatch repair protein hexB] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:g1002521] [LN:BSU27343] [AC:U27343] [PN:MutL] [GN:mutL] [FN:mismatch-repair recognition] [OR:Bacillus subtilis] [DB:genpept-bctl] [DE:Bacillus subtilis spore coat protein (cotE) gene, partial cds, andmismatch repair recognition proteins (mutS) and (mutL) genes, complete cds.] [LE:2782] [RE:4665] [DI:direct] >gp:[GI:e1185296:g2634077] [LN:BSUB0009] [AC:Z99112:AL009126] [GN:mutL] [FN:DNA mismatch repair] [OR:Bacillus subtilis] [DB:genpept-bctl] [DE:Bacillus subtilis complete genome (section 9 of 21): from 1598421to 1807200.] [SP:P49850] [LE:179285] [RE:181168] [DI:direct]

4300

 ORF Name
 NTID
 AAID
 NT AA Length Length
 Score
 probability

 A17503000983_26306257_c2_835
 529
 4301
 2073
 690
 2490
 1.0e-258

Description

sp:[LN:TOP1_BACSU] [AC:P39814] [GN:TOPA:TOPI] [OR:BACILLUS SUBTILIS] [EC:5.99.1.2]
[DE:(UNTWISTING ENZYME) (SWIVELASE)] [SP:P39814] [DB:swissprot] >pir:[LN:G69724]
[AC:G69724] [PN:DNA topoisomerase I topA] [GN:topA] [CL:DNA topoisomerase I]
[OR:Bacillus subtilis] [DB:pir2] >gp:[GI:g520753] [LN:BACSMF] [AC:L27797] [PN:DNA
topoisomerase I] [FN:DNA unwinding protein: removes negative] [OR:Bacillus subtilis]
[SR:Bacillus subtilis (strain 8G5) DNA] [DB:genpept-bct1] [DE:Bacillus subtilis (smf)
gene, 3' end, DNA topisomase gene, completecds, (gid) gene, 5' end.] [LE:673] [RE:2748]
[DI:direct] >gp:[GI:e1185203:g2633984] [LN:BSUB0009] [AC:Z99112:AL009126] [PN:DNA
topoisomerase I] [GN:topA] [FN:DNA unwinding protein removing negative] [OR:Bacillus
subtilis] [DB:genpept-bct1] [EC:5.99.1.2] [DE:Bacillus subtilis complete genome
(section 9 of 21): from 1598421to 1807200.] [NT:alternate gene name: topI] [SP:P39814]
[LE:84640] [RE:86715] [DI:direct] >gp:[GI:e332180:g2462970] [LN:BSYLQGCOD]
[AC:AJ000975] [PN:DNA Topoisomerase I] [GN:topA] [OR:Bacillus subtilis]
[DB:genpept-bct1] [DE:Bacillus subtilis ylqg to codV gene region.] [SP:P39814]
[LE:3808] [RE:5883] [DI:direct]

 ORF Name
 NTID
 AAID
 NT AA Length Length
 Score
 probability

 A17503000983_26353417_c1_753
 530
 4302
 294
 97
 141
 8.5e-10

Description

sp:[LN:YVI2_CLOPE] [AC:Q46213] [OR:CLOSTRIDIUM PERFRINGENS] [DE:HYPOTHETICAL 10.7 KD PROTEIN IN VIRR 5'REGION (ORF2)] [SP:Q46213] [DB:swissprot] >pir:[LN:S49553] [AC:S49553] [PN:hypothetical protein 2] [OR:Clostridium perfringens] [DB:pir2] >gp:[GI:g498839] [LN:CPVIRRS] [AC:U04966] [FN:unknown] [OR:Clostridium perfringens] [DB:genpept-bct1] [DE:Clostridium perfringens JIR4025 extracellular toxin productionregulatory locus ORF1 and ORF3 genes, partial cds, and ORF2,ORF10c, virR, virS, and ORF4 genes, complete cds.] [NT:ORF2] [LE:469] [RE:756] [DI:direct]

ORF Name	NTID AAID LengthLength score probability
AI7503000983_26354837_c2_946	531 4303 900 299 570 2.9e-55
[EC:4.2.1.52] [DE:DIHYDRODIPICOLINAT >pir:[LN:E64330] [AC:E64330] [PN:di jannaschii] [EC:4.2.1.52] [DB:pir2] [LN:U67480] [AC:U67480:L77117] [PN:d [OR:Methanococcus jannaschii] [DB:ge	DAPA:MJ0244] [OR:METHANOCOCCUS JANNASCHII] TE SYNTHASE, (DHDPS)] [SP:Q57695] [DB:swissprot] ihydrodipicolinate synthase,] [OR:Methanococcus [MP:REV233451-232582] >gp:[GI:g1590977] dihydrodipicolinate synthase (dapA)] [GN:MJ0244] enpept-bct2] [DE:Methanococcus jannaschii section 22 :similar to GB:L08471 SP:Q04796 PID:142830] [LE:511]
ORF Name	NTID AAID NT AA score probability
A17503000983_26369016_c1_773	532 4304 387 128 105 5.6e-06
	[AC:X84706] [PN:scaffolding protein] [GN:bl hpept-phg] [DE:Bacteriophage TP901-1 genomic [358] [DI:direct]
ORF Name	NTID AAID NT AA score probability
A17503000983_26460951_c2_839	533 4305 792 263 917 5.0e-92
<pre>biosynthesis] [OR:Bacillus subtilis]</pre>	[PN:uridylate kinase] [GN:smbA] [FN:pyrimidine [DB:genpept-bct1] [EC:2.7.4] [DE:Bacillus of 21): from 1598421to 1807200.] [LE:120774]
ORF Name AI7503000983_26569432_f2_444	NTID AAID NT AA score probability S34
Description NO-HIT	
ORF Name	NTID AAID NT AA score probability
AI7503000983_26586537_f2_376	535 4307 156 51 70 0.028
<pre>protein] [GN:APE2412] [OR:Aeropyrum]</pre>	pernix] [SR:Aeropyrum pernix (strain:K1) DNA] enomic DNA, section 7/7.] [LE:105148] [RE:105525]
ORF Name	NTID AAID NT AA score probability
A17503000983_2734778_f1_31	536 4308 126 41
Description NO-HIT	

ORF Name	NTID	AAID	NT AA LengthLength score probability
AI7503000983_275427_c1_744	537	4309	1379 5.5e-141
Description pir:[LN:H69884] [AC:H69884] [PN:con [CL:conserved hypothetical protein M >gp:[GI:e1185269:g2634050] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-b 9 of 21): from 1598421to 1807200.] [RE:152056] [DI:direct]	G139] [9 009] [A ct1] [D	OR:Bac C:Z991 E:Baci	cillus subtilis] [DB:pir2] 112:AL009126] [GN:ymfA] [FN:unknown] illus subtilis complete genome (section
ORF Name	NTID	AAID	NT AA LengthLength score probability
AI7503000983_2931337_f2_256	538	4310	780 259 771 1.5e-76
gp:[GI:g3800827] [LN:AF076684] [AC:A ATPase domain] [GN:opp-2D] [OR:Staph; [DE:Staphylococcus aureus oligopeption (opp-2B), oligopeptide transporter proligopeptide transporter coligopeptide transporterputative ATPase domain (of [DI:direct]	ylococc de trans utatives ase doma	us aur sporte membra ain (o	reus] [DB:genpept-bct2] er putative membranepermease domain ane permease domain (opp-2C), opp-2D), and oligopeptide
ORF Name	NTID	AAID	NT AA LengthLength score probability
AI7503000983_29781968_c3_995	539	4311	855 284 825 2.8e-82
>gp:[GI:g142997] [LN:BACGLPPFK] [AC:I [OR:Bacillus subtilis] [SR:Bacillus subtilis antiterminator regulatory progenes, complete cds, glycerolkinase [RE:1909] [DI:direct] >gp:[GI:e11829] [PN:glycerol uptake facilitator] [GN subtilis] [DB:genpept-bct1] [DE:Bacilfrom 802821 to1011250.] [SP:P18156] >gp:[GI:e1182929:g2633263] [LN:BSUB06] facilitator] [GN:glpF] [FN:glycerol uptake facillus subtilitator] [DB:genpept-bct1] [DE:Bacillus subtilitator] [CR:BSY14079] [AC:Y14079] [PN:putative [CR:Bacillus subtilis] [DB:genpept-bct]	swissp: N:glyce: R:Bacil: M99611] Subtilia rotein (glpK) (17:g2633 :glpF] llus sul [LE:1995 006] [Ac utilizat lis comp E:3330] We glyce ct1] [Di ceam.]	rot] > rot] > rol up lus su [PN:gs DNA] (glpP) gene, 3251] [FN:gl btilis 186] [C:Z991 tion] plete [DI:derol u E:Baci	pir:[LN:C47700] ptake facilitator glpF] [GN:glpF] pubtilis] [DB:pir2] [MP:75 (degrees)] glycerol uptake facilitator] [GN:glpF] [DB:genpept-bct1] [DE:Bacillus [Oglycerol uptake facilitator (glpF) [S' end.] [NT:putative] [LE:1085] [LN:BSUB0005] [AC:Z99108:AL009126] [lycerol utilization] [OR:Bacillus [Complete genome (section 5 of 21): [RE:200010] [DI:direct] [O9:AL009126] [PN:glycerol uptake [OR:Bacillus subtilis] [OR:Bacillus subtilis] [OR:GI:e324940:g2226136]
ORF Name	NTID	AAID	NT AA LengthLength score probability

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000983_30355313_c1_779	541	4313	1113	370	382	2.5e-35
Description		L			J	
pir:[LN:C69901] [AC:C69901] [PN:pro [GN:yocF] [CL:probable Bacillus sub [OR:Bacillus subtilis] [DB:pir2] >gp [PN:sensor kinase] [GN:yocF] [OR:Bac subtilis chromosome region between t sensor kinase (385 aa)] [LE:70568] [LN:BSUB0011] [AC:Z99114:AL009126] [DB:genpept-bct1] [DE:Bacillus subti 2000171to 2207900.] [NT:similar to t [RE:90771] [DI:direct]	tilis to e:[GI:g26 illus so erC and RE:71680 GN:yocF] lis comp	wo-comp 619013] ubtilis odhAB. 0] [DI:] [FN:u	[LN:A [LN:A] [DB:] [NT: direct unknown	sensor F02786 genper simila] >gp: [OR: (secti	hist: [8] [AG ot-bct] ar to I [GI:e] Bacill on 11	idine kinase yocF] C:AF027868] L] [DE:Bacillus B.subtilis DegS L185391:g2634312] Lus subtilis] of 21): from
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000983_30656317_c2_838	542	4314	822	273	858	8.9e-86
Description						
pir:[LN:S61496] [AC:S61496:H69601] [GN:codY] [OR:Bacillus subtilis] [D [PN:CodY] [GN:codY] [OR:Bacillus sub JH642 dipeptide permease operon regucds.] [LE:3225] [RE:4004] [DI:direct [AC:Z99112:AL009126] [PN:transcripti of srfA and comK genes (in] [OR:Baci subtilis complete genome (section 9 [LE:91098] [RE:91877] [DI:direct]	B:pir2] tilis] lators,] >gp:[0 onal rec llus sub	>gp:[G [DB:gen codV,c GI:e118 gulator ptilis]	I:g535 pept-b odW, c 5208:g [GN: DB:g	351] [ct1] [cdX, accordY] codY]	LN:BSU DE:Bac nd coc 9] [LN [FN:ne -bct1]	J13634] [AC:U13634] cillus subtilis dY genes, complete N:BSUB0009] egative regulation [DE:Bacillus
ORF Name	NTID	<u>AAID</u>	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000983_31275_c2_927	543	4315	171	56	;	
Description NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000983_31552_c1_755	544	4316	522	173	197	9.9e-16
Description gp:[GI:d1045212:g5106115] [LN:AP0000 protein] [GN:APE2411] [OR:Aeropyrum [DB:genpept] [DE:Aeropyrum pernix ge: coupled receptors signature] [LE:104	pernix] nomic DN	[SR:Ae NA, sec	ropyru tion 7	m pern /7.] [ix (st NT:mot	rain:K1) DNA] if=G-protein
ORF Name	NTID	AAID	<u>NT</u> Lengthl	<u>AA</u> Length	score	probability
A17503000983_31697151_c2_826	545	4317	315	104	441	1.4e-41
Description						
sp:[LN:RL19_BACSU] [AC:O31742] [GN:R PROTEIN L19] [SP:O31742] [DB:swisspre protein L19 rplS] [GN:rplS] [CL:Esc subtilis] [DB:pir2] >gp:[GI:e1185195 [PN:ribosomal protein L19] [GN:rplS] [DE:Bacillus subtilis complete genome [SP:O31742] [LE:77012] [RE:77368] [D	ot] >pir herichia :g263397 [OR:Bac e (secti	::[LN:E coli [6] [LN cillus on 9 o	69696] riboso :BSUBO subtil:	AC:E mal pr 009] [is] [D	69696 otein AC:Z99 B:genp] [PN:ribosomal L19] [OR:Bacillus 112:AL009126] ept-bct1]

ORF Name	NTID	AAID LengthLength score probability
AI7503000983_31803760_£1_9	546	4318 153 50 43 0.031
Description		
	rhabdit	Z81556] [GN:F58G1.9] [OR:Caenorhabditis is elegans cosmid F58G1, complete sequence.] 3426:13549] [RE:13379:13500:13656]
ORF Name	NTID	AAID NT AA probability
A17503000983_31844658_c3_987	547	4319 138 45
Description		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000983_3235828_c1_798	548	4320 138 45
Description		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000983_3257827_c2_858	549	4321 705 234 356 1.4e-32
Description		
[DB:pir2] >gp:[GI:e1185278:g2634059]	[LN:BS] [DB:gen] 21to 180	
ORF Name	NTID	AAID NT AA score probability
A17503000983_32611557_c3_966	550	4322 294 97 316 2.4e-28
Description sp:[LN:EFTS_BACSU] [AC:P80700:O31748 FACTOR TS (EF-TS)] [SP:P80700:O31748		SF] [OR:BACILLUS SUBTILIS] [DE:ELONGATION wissprot]
ORF Name	NTID	AAID NT AA score probability
A17503000983_33283167_c3_983	551	4323 1338 445 967 2.5e-97
Description		
	:[GI:e11 nknown] e (secti	L85277:g2634058] [LN:BSUB0009] [OR:Bacillus subtilis] [DB:genpept-bct1] ion 9 of 21): from 1598421to 1807200.]
ORF Name	NTID	AAID LengthLength score probability
AI7503000983_33375260_t2_273	552	4324 165 54
Description NO-HIT		
740-11TT		

ORF Name AI7503000983_3394390_f1_48	<u>NTID</u>	AAID NT AA score probability 4325 123 40
Description NO-HIT		
ORF Name A17503000983_34070261_c3_1010 Description pir: [LN:A70039] [AC:A70039] [PN:ABC [GN:yvfR] [CL:ATP-binding cassette >gp: [GI:e1186097:g2635922] [LN:BSUB00] [OR:Bacillus subtilis] [DB:genpept-books of 21): from 3399551to 3609060.] protein)] [LE:97865] [RE:98770] [DI:0] [AC:Z94043] [PN:hypothetical protein)	homology 018] [AC ctl] [DE [NT:simi compleme] [GN:yv omic DNA	C:Z99121:AL009126] [GN:yvfR] [FN:unknown] E:Bacillus subtilis complete genome (section ilar to ABC transporter (ATP-binding ent] >gp:[GI:e313073:g1945718] [LN:BSZ94043] vfR] [OR:Bacillus subtilis] A fragment (88 kb).] [NT:probable ABC
dehydratase] [OR:Bacillus subtilis] [LN:BSTHRBC] [AC:X04603] [PN:threoning [DB:genpept-bct1] [EC:4.2.99.2] [DE:Rkinase and threoninesynthase (EC 2.7 [LE:248] [RE:1306] [DI:direct] >gp:[CAC:Z99120:AL009126] [PN:threonine symplements of the control	[DB:swinine synferm sy	issprot] >pir:[LN:A25364] inthase, thrC] [GN:thrC] [CL:threonine .99.2] [DB:pir2] >gp:[GI:g40211] inase] [GN:thrC] [OR:Bacillus subtilis] iilis thrB and thrC genes for homoserine and EC 4.2.99.2, respectively).] [SP:P04990] 4304:g2635722] [LN:BSUB0017] [[GN:thrC] [FN:threonine biosynthesis] C:4.2.99.2] [DE:Bacillus subtilis complete 414420.] [NT:alternate gene name: thrB]
ORF Name AI7503000983_34257817_c3_1038 Description NO-HIT	<u>NTID</u> 556	AAID NT AA score probability 4328 123 40
ORF Name AI7503000983_34571877_c1_746 Description gp:[GI:g3426364] [LN:AF082738] [AC:AF [DB:genpept-bct2] [DE:Streptococcus p		AAID NT AA LengthLength score probability 4329 1275 424 484 3.8e-46 [PN:unknown] [OR:Streptococcus pyogenes] sphosphotidylglycerophosphate synthase

(pgsA) and ABC transporter ATP-binding protein (stpA) genes, complete cds; and unknown genes.] [NT:orf1] [LE:1] [RE:1245] [DI:direct]

ORF Name	NTID	AAID.	<u>NT</u> Length	AA Length sco	re pr	obability	
A17503000983_34663177_c2_860	558	4330	1167	388 641	2.1e	-65	
Description gp:[GI:g1842440] [LN:BSU87792] [AC:U competence-damage inducible function [DE:Bacillus subtilis tRNA-Ala, phos (cinA) genes, complete cds, and RecA [DI:direct]] [OR:B phatidy	acillu: lglyce:	s subti rophosp	lis] [DB:q hate synt]	genpept nase (pg	-bct1] sA) and CinA	
ORF Name	NTID	AAID	<u>NT</u> Lengthl	AA Length sco	re pr	obability	
A17503000983_35187587_c3_1055	559	4331		195 370	4.6e	-34	
Description pir:[LN:S34747] [AC:S34747] [PN:ant amidotransferase:trpG homology] [OR:							
ORF Name	NTID	AAID	NT Length1	AA Length sco	ce pr	obability	
A17503000983_35370318_c2_834	560	4332	1930	309 127	4 7.4e	-130	
Description			J				
pir:[LN:F69719] [AC:F69719] [PN:suc [GN:sucD] [CL:succinateCoA ligase [EC:6.2.1.5] [DB:pir2] >gp:[GI:e1185] [PN:succinyl-CoA synthetase (alpha s [DB:genpept-bct1] [EC:6.2.1.5] [DE:B from 1598421to 1807200.] [SP:P80865] >gp:[GI:e332178:g2462968] [LN:BSYLQG:synthetase alpha chain] [GN:sucD] [O:subtilis ylqg to codV gene region.]	(ADP-f 201:g26 ubunit) acillus [LE:82 COD] [A R:Bacil	orming) 33982]] [GN:: subtil 596] [I C:AJ000 lus sul	alpha [LN:BS] sucD] [0 lis comp RE:83498 0975] [1 otilis]	chain] [0 DB0009] [1 DR:Bacillu plete geno B] [DI:din PN:putativ [DB:genpe	DR:Baci AC:Z991 us subt ome (se rect] re succ ept-bct	llus subtilis] 12:AL009126] ilis] ction 9 of 21) inyl-coA 1] [DE:Bacillus	
ORF Name	NTID	AAID	<u>NT</u> LengthI	AA Length	re pro	obability	
A17503000983_35557787_c3_1073	561	4333		424 842		-84	
Description		<u> </u>	J				
gp:[GI:g4982084] [LN:AE001799] [AC:A [OR:Thermotoga maritima] [DB:genpept of the complete genome.] [NT:similar [LE:15047] [RE:16252] [DI:complement	-bct2] to PID	[DE:The	ermotoga	a maritima	secti	on 111 of 136]
ORF Name	NTID	AAID	<u>NT</u> LengthI	AA ength	re pro	obability	_
AI7503000983_36134401_c2_897	562	4334	831	276 315	3.1e	-28	
Description							
sp:[LN:YXEH_BACSU] [AC:P54947] [GN:Y30.2 KD PROTEIN IN IDH-DEOR INTERGENDED Spir:[LN:B70075] [AC:B70075] [PN:COME [CL:Methanobacterium thermoautotroph:[OR:Bacillus subtilis] [DB:pir2] >gp [AC:Z99124:AL009126] [GN:yxeH] [FN:um [DE:Bacillus subtilis complete genome [NT:similar to hypothetical proteins >gp:[GI:d1008920:g1408493] [LN:D45913] [SR:Bacillus subtilis (strain:BGSC 120 [DE:Bacillus subtilis genome sequence complete cds.] [NT:homologous to Swise	IC REGIONS REG	ON] [SI hypoth nserved 184680: [OR:Ba ion 21 54947] D45912] burg 16	e:P5494 netical hypoth g263650 acillus of 21): [LE:633 [GN:y2 58; trp0 iol and	7] [DB:swi protein y netical pr 01] [LN:BS subtilis] from 399 .94] [RE:6 ceH] [OR:E C2)) DNA]	ssprot xeH] [0 otein 1 UB0021 [DB:go 9281to 4006] acillus [DB:go on,par] GN:yxeH] MTH1071]] enpept-bct1] 4214814.] [DI:complement] s subtilis] npept-bct1] tial and	

ORF Name	NTID	AAID	<u>NT</u> LengthL	<u>AA</u> ength ^S	core	probability	
A17503000983_36142827_c1_814	563	4335	1812 6	03	780 1	6e-77	
Description gp:[GI:d1014255:g1651216] [LN:D88209 licheniformis] [SR:Bacillus lichenifo [DE:Bacillus licheniformis DNA for P	ormis (strain	:N22) DN	A] [DB	:genpe	ept-bct1]	
ORF Name AI7503000983 36220061 c3 964	NTID	AAID [4336	<u>NT</u> LengthL			probability	
Description			ا لـــــال		-الــــــــــــــــــــــــــــــــــــ		
sp:[LN:HSLU_BACSU] [AC:P39778] [GN:HSSHOCK PROTEIN HSLU] [SP:P39778] [DB:EAC:E69601:S61495:S72310] [PN:ATP-deported protein] [GN:clpY:codX] [CL:heat shous and showed protein] [GN:clpY:codX] [GN:codX] [GN:CdX] [GN:CdX] [GN:DdX] [GX:Z99112:AL009126] [PN:ATP-dependent subtilis] [DB:genpept-bct1] [DE:Bacin from 1598421to 1807200.] [NT:alternation [RE:91058] [DI:direct]	swisspropered and several seve	ot] >p: t Clp p tein h :pir2] s subt: gulator ct] >gp proteas otilis	ir:[LN:E proteina slU:FtsH >gp:[GI ilis] [D cs, codV p:[GI:e1 se-like] complet	69601] se-like /SEC18 :g5353 B:genpe ,codw, 185207 [GN:c] e genoe	e prot /CDC48 50] [I ept-bc codX, :g2633 lpY] [cein clpY:codX 3-type ATP-bindin LN:BSU13634] ct1] [DE:Bacillus and codY genes, 3988] [LN:BSUB000 [OR:Bacillus ection 9 of 21):	
ORF Name	NTID	AAID	<u>NT</u> LengthLe	AA ength s	core	probability	w 464
AI7503000983_36225938_c2_833	565	4337][1185][3	94 1	557 7	.6e-160	
Description							
sp:[LN:SUCC_BACSU] [AC:P80886] [GN:SI [DE:(VEGETATIVE PROTEIN 63) (VEG63)] [AC:E69719] [PN:succinateCoA ligas [CL:succinateCoA ligase (ADP-forming [DB:pir2] >gp:[GI:e1185200:g2633981] [PN:succinyl-CoA synthetase (beta substitute [DB:genpept-bct1] [EC:6.2.1.5] [DE:Bafrom 1598421to 1807200.] [SP:P80886] >gp:[GI:e332177:g2462967] [LN:BSYLQGO synthetase beta chain] [GN:sucC] [OR subtilis ylqg to codV gene region.]	[SP:P86 se (ADP- ng) beta [LN:BS0 ounit)] acillus [LE:814 COD] [A0 :Bacillu	D886] -forming chair (JB0009] [GN:susubtil 410] [FC:AJ000	[DB:swis ng), bet n] [OR:B [AC:Z9 ncC] [OR lis comp RE:82567 0975] [Pi	sprot] a chain acillum 9112:Al :Bacill lete ge] [DI:c N:putan DB:gen	>pir: n] [GN s subt L00912 lus su enome direct tive s pept-b	[LN:E69719] U:sucC] Cilis] [EC:6.2.1. C6] Cbtilis] (section 9 of 21 C] Succinyl-coA Cct1] [DE:Bacillu):
ORF Name	NTID	AAID	<u>NT</u> LengthLe	AA ength	core	probability	
A17503000983_3913307_±3_640	566	4338	<u> </u>				
Description		L					
NO-HIT							
ORF Name	NTID	AAID	<u>NT</u> LengthLe	AA ength	core	probability	
A17503000983_3928177_f2_255	567	4339	360 1	19 9	1 0	.00081	
Description							
<pre>gp:[GI:d1039115:g4514335] [LN:AB01337 halodurans] [SR:Bacillus halodurans halodurans C-125 yndF, gerKA, yndFl a [LE:1025] [RE:1876] [DI:direct]</pre>	(strain:	C-125)	DNA] [1	DB:genr	pept-b	ctl] [DE:Bacillus	

ORF Name	NTID	AAID	<u>NT</u> LengthLe	AA ength sco	re probab	ility
A17503000983_3937551_c1_820	568	4340		32 803	6.0e-80	
Description		L	<u> </u>			
pir:[LN:B70461] [AC:B70461] [PN:asp [CL:aspartate-semialdehyde dehydroge >gp:[GI:g2984139] [LN:AE000760] [AC: dehydrogenase] [GN:asd] [OR:Aquifex section 92 of 109 of the complete ge	nase] [C AE000760 aeolicus	OR:Aqui O:AE000 3] [DB:	fex aeo 0657] [Pl genpept	licus] [] N:asparta -bct2] []	EC:1.2.1.11 ate-semiald DE:Aquifex] [DB:pir2] ehyde
ORF Name	NTID	AAID	<u>NT</u> LengthLe	AA ength	re probab	ility
A17503000983_3961702_c1_811	569	4341	1269 4	22 224	9 3.5e-233	
<pre>Description pir:[LN:JC5325] [AC:JC5325:PC4317] [CL:methicillin resistance factor fe</pre>						-
ORF Name	NTID	AAID	<u>NT</u> LengthLe	AA ength sco	re probab	ility
AI7503000983_4025303_c2_923	570	4342	792 20	63 450	1.5e-42	
sp:[LN:TRPC_LACLA] [AC:Q01999] [GN:T [SR:,SUBSPLACTIS:STREPTOCOCCUS LACTI SYNTHASE, (IGPS)] [SP:Q01999] [DB:sw [PN:indole-3-glycerol-phosphate synt synthase:trpC homology] [OR:Lactococ >gp:[GI:g149519] [LN:LACTRPOP] [AC:M [GN:trpC] [OR:Lactococcus lactis] [S lactis) DNA] [DB:genpept-bct1] [EC:4 trpC, trpB trpA genes, completecds.]	S] [EC:4 rissprot] hase,] [cus lact [87483] [R:Lactoc 1.1.48]	>pir: Spir: GN:trp Spir: Spir:	[LN:S35] [CLN:S35] [CL] [Sp. lact cloleglyce lactis [] [] [] [] [] [] [] [] [] [] [] [] []	INDOLE-3- 127] [AC :indole-1 tis] [EC erol phos (strain 1 s trpE, t	:S35127] 3-glycerol- :4.1.1.48] sphate synt IL1403, sub	phosphate [DB:pir2] hase] _species
ORF Name	NTID	AAID	<u>NT</u> Length Le	AA ength sco	re probab	ility
A17503000983_4062762_£3_555	571	4343		34 508	1.1e-48	
Description gp:[GI:g3135292] [LN:AF029731] [AC:A channel] [GN:mscL] [OR:Staphylococcu aureus large conductance mechanosens [LE:49] [RE:411] [DI:direct]	s aureus	[DB:	genpept -	-bct2] [I	E:Staphylo	coccus
ORF Name	NTID	AAID	<u>NT</u> LengthLe	AA ength scor	re probab	ility
AI7503000983_4063802_c1_808	572	4344	1497 43	98 808	1.8e-80	
Description sp:[LN:TRPE_LACLA] [AC:Q02001] [GN:T [SR:,SUBSPLACTIS:STREPTOCOCCUS LACTI COMPONENT I,] [SP:Q02001] [DB:swissp synthase, alpha chain] [GN:trpE] [C [OR:Lactococcus lactis subsp. lactis [LN:LACTRPOP] [AC:M87483] [PN:anthra [OR:Lactococcus lactis] [SR:Lactococ [DB:genpept-bct1] [EC:4.1.3.27] [DE:	S] [EC:4 rot] >pi L:anthra] [EC:4. nilate s cus lact	.1.3.2 r:[LN: nilate 1.3.27 ynthas is (st	7] [DE:F S35124] synthas] [DB:pi e alpha rain IL1	ANTHRANII [AC:S351 se compor ir2] >gp: subunit] 1403, sub	.24] [PN:an ment I] [GI:g149510 [GN:trpE] D_species la	nthranilate 6] actis) DNA]

genes, completecds.] [LE:954] [RE:2324] [DI:direct]

ORF Name	NTID	AAID LengthLength score probability
A17503000983_40686_c1_737	573	4345 4089 1362 6425 0.0
[DE:DNA POLYMERASE III, ALPHA CHAIN [DB:swissprot] >gp:[GI:d1013849:g148 polymerase III] [OR:Staphylococcus a	POLC-TY 3182] [ureus] Staphyl	LN:D86727] [AC:D86727:D45368] [PN:DNA [SR:Staphylococcus aureus (strain:IP8) DNA, ococcus aureus DNA for DNA polymerase III,
ORF Name	NTID	AAID NT AA probability
A17503000983_4069675_£2_274	574	4346 162 53
Description		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000983_4080342_c3_996	575	4347 1515 504 1998 1.4e-206
Sp:[GI:g142992] [LN:BACGLPKD] [AC:M (strain W168) DNA] [DB:genpept-bct1] glycerol-3-phosphatedehydrogenase (g (glpK) (EC 2.7.1.30)] [LE:698] [RE:2 [LN:BSUB0005] [AC:Z99108:AL009126] [atilization] [OR:Bacillus subtilis] subtilis complete genome (section 5 [LE:200029] [RE:201519] [DI:direct] [AC:Z99109:AL009126] [PN:glycerol ki [OR:Bacillus subtilis] [DB:genpept-b genome (section 6 of 21): from 99950 [DI:direct] >gp:[GI:e324941:g2226137 [GN:glpK] [OR:Bacillus subtilis] [DB] [DB:s N:glyce lis] [E 34393] [DE:B. lpD) ge 188] [D PN:glyc [DB:gen of 21): >gp:[GI nase] [ct1] [E 1 to120] [LN:B :genpep n and d	wissprot] >pir:[LN:B45868] rol kinase, glpK] [GN:glpK] C:2.7.1.30] [DB:pir2] [MP:75 (degrees)] [OR:Bacillus subtilis] [SR:B.subtilis subtilis glycerol kinase (glpK) and nes, complete cds.] [NT:glycerol kinase I:direct] >gp:[GI:e1182918:g2633252] erol kinase] [GN:glpK] [FN:glycerol pept-bct1] [EC:2.7.1.30] [DE:Bacillus from 802821 to1011250.] [SP:P18157] :e1182930:g2633264] [LN:BSUB0006] GN:glpK] [FN:glycerol utilization] C:2.7.1.30] [DE:Bacillus subtilis complete 9940.] [SP:P18157] [LE:3349] [RE:4839] SY14079] [AC:Y14079] [PN:glycerol kinase] t-bct1] [DE:Bacillus subtilis chromosomal ownstream.] [NT:see EMBL M34393 and Swiss
DRF Name A17503000983_4089062 c2 906	NTID	AAID NT AA score probability
Description	576	4348 1176 391 380 4.0e-35
pir:[LN:C71302] [AC:C71302] [PN:pro	ilis sp	xonuclease] [GN:TP0626] [OR:Treponema irochete] [DB:pir2] >gp:[GI:g3322921] onuclease. putative] [GN:TP0626]

[OR:Treponema pallidum] [DB:genpept-bct2] [DE:Treponema pallidum section 53 of 87 of the complete genome.] [NT:similar to SP:P23479 percent identity: 32.68;] [LE:11246] [RE:12421] [DI:direct]

ORF Name	NTID	AAID LengthLength score probability
A17503000983_4093818_c2_859	577	4349 591 196 436 4.7e-41
Description		
sp:[LN:PGSA_BACSU] [AC:P46322] [GN:F	egsal [or	R:BACILLUS SUBTILIS] [EC:2.7.8.5] [DE:(EC
2.7.8.5) (PHOSPHATIDYLGLYCEROPHOSPHA		
[DB:swissprot] >gp:[GI:d1009402:g893		
_		s (strain:Marburg168) DNA] [DB:genpept-bct1]
		ne for phosphatidylglycerophosphatesynthase,
complete cds.] [LE:182] [RE:763] [DI		
		e synthase] [GN:pgsA] [FN:involved in the
_	_	subtilis] [DB:genpept-bct1] [DE:Bacillus
		hate synthase(pgsA) and CinA (cinA) genes,
complete cds, and RecA (recA) gene, p		
[DI:direct]		odo:, [gan, [, [
ORF Name	NTID	AAID NT AA score probability
		LengthLength
A17503000983_4095286_c2_869	578	4350 375 124 246 6.4e-21
Description		
pir:[LN:C69884] [AC:C69884] [PN:con	served h	hypothetical protein ymcA] [GN:ymcA]
[OR:Bacillus subtilis] [DB:pir2] >gr	:[GI:e11	185293:g2634074] [LN:BSUB0009]
[AC:Z99112:AL009126] [GN:ymcA] [FN:u	nknown]	[OR:Bacillus subtilis] [DB:genpept-bct1]
[DE:Bacillus subtilis complete genom	e (secti	ion 9 of 21): from 1598421to 1807200.]
[NT:similar to hypothetical proteins	[LE:17	75322] [RE:175753] [DI:direct]
		NTI 22
ORF Name	NTID	AAID NT AA score probability
ORF Name		LengthLength score probability
ORF Name AI7503000983_4334383_f2_322	<u>NTID</u> 579	AATD — geore probability
ORF Name AI7503000983_4334383_f2_322 Description		LengthLength score probability
ORF Name AI7503000983_4334383_f2_322		LengthLength score probability
ORF Name A17503000983_4334383_f2_322 Description NO-HIT	579	AAID LengthLength score probability 4351 135 44 AAID NT AA ggore probability
ORF Name AI7503000983_4334383_f2_322 Description		AAID LengthLength score probability 4351 135 44
ORF Name A17503000983_4334383_f2_322 Description NO-HIT	579	AAID LengthLength score probability 4351 135 44 AAID NT AA ggore probability
ORF Name A17503000983_4334383_f2_322 Description NO-HIT ORF Name	NTID	AAID LengthLength score probability 4351 135 44 AAID NT AA LengthLength score probability
ORF Name AI7503000983_4334383_f2_322 Description NO-HIT ORF Name AI7503000983_4336536_c1_739 Description	579 NTID 580	AAID LengthLength score probability AAID NT AA LengthLength score probability LengthLength 211 3.3e-17
ORF Name AI7503000983_4334383_f2_322 Description NO-HIT ORF Name AI7503000983_4336536_c1_739 Description	NTID [580] [LXR] [OR	AAID LengthLength score probability AAID NT AA LengthLength score probability AAID LengthLength score probability A352 315 104 211 3.3e-17 R:BACILLUS SUBTILIS] [DE:HYPOTHETICAL 10.4
ORF Name A17503000983_4334383_f2_322 Description NO-HIT ORF Name A17503000983_4336536_c1_739 Description sp:[LN:YLXR_BACSU] [AC:P32728] [GN:YKD PROTEIN IN NUSA-INFB INTERGENIC R	NTID [580] LXR] [OF EGION (C	AAID LengthLength AAID NT AA LengthLength score probability AAID LengthLength score probability 4352 315 104 211 3.3e-17 R:BACILLUS SUBTILIS] [DE:HYPOTHETICAL 10.4 DRF3)] [SP:P32728] [DB:swissprot]
ORF Name A17503000983_4334383_f2_322 Description NO-HIT ORF Name A17503000983_4336536_c1_739 Description sp:[LN:YLXR_BACSU] [AC:P32728] [GN:YLXR_BACSU] [AC:P32728] [GN:YLXR_BACSU] [AC:P32728] [GN:YLXR_BACSU] [AC:D36905:D69882:S	NTID [580 LXR] [OF EGION (C 31992]	AAID LengthLength score probability AAID NT AA LengthLength score probability AAID LengthLength score probability A352 315 104 211 3.3e-17 R:BACILLUS SUBTILIS] [DE:HYPOTHETICAL 10.4 DRF3)] [SP:P32728] [DB:swissprot] [PN:conserved hypothetical protein
ORF Name AI7503000983_4334383_f2_322 Description NO-HIT ORF Name AI7503000983_4336536_c1_739 Description sp:[LN:YLXR_BACSU] [AC:P32728] [GN:YLXR_BACSU] [AC:P32728] [GN:YLXR_BACSU] [AC:P32728] [GN:YLXR_BACSU] [AC:D36905:D69882:SylxR:hypothetical protein 1 (nusA 3')	NTID [580] LXR] [OR EGION (C) 31992] region)	AAID LengthLength AAID NT AA LengthLength score probability AAID LengthLength score probability 4352 315 104 211 3.3e-17 R:BACILLUS SUBTILIS] [DE:HYPOTHETICAL 10.4 DRF3)] [SP:P32728] [DB:swissprot] [PN:conserved hypothetical protein [] [GN:ylxR] [OR:Bacillus subtilis]
ORF Name A17503000983_4334383_f2_322 Description NO-HIT ORF Name A17503000983_4336536_c1_739 Description sp:[LN:YLXR_BACSU] [AC:P32728] [GN:YLXR_BACSU] [AC:P32728] [GN:YLXR_BACSU] [AC:D36905:D69882:SylxR:hypothetical protein 1 (nusA 3'[DB:pir2] >gp:[GI:g580900] [LN:BSORF	NTID [580 LXR] [OR EGION (C 31992] region) 1T7A] [A	AAID LengthLength AAID LengthLength AAID LengthLength LengthLength Score probability AAID LengthLength LengthLength LengthLength Score probability AC:BACILLUS SUBTILIS] [DE:HYPOTHETICAL 10.4 DRF3)] [SP:P32728] [DB:swissprot] [PN:conserved hypothetical protein [In:Sp:P32728] [OR:Bacillus subtilis] AC:Z18631] [GN:ORF3] [OR:Bacillus subtilis]
ORF Name AI7503000983_4334383_f2_322 Description NO-HIT ORF Name AI7503000983_4336536_c1_739 Description sp:[LN:YLXR_BACSU] [AC:P32728] [GN:YKD PROTEIN IN NUSA-INFB INTERGENIC R >pir:[LN:D36905] [AC:D36905:D69882:S ylxR:hypothetical protein 1 (nusA 3' [DB:pir2] >gp:[GI:g580900] [LN:BSORF [DB:genpept-bct1] [DE:B.subtilis inf	NTID [580 LXR] [OR EGION (C 31992] region) 1T7A] [A B-nusA c	AAID LengthLength AAID LengthLength AAID LengthLength LengthLength Score probability AAID LengthLength LengthLength 211 3.3e-17 R:BACILLUS SUBTILIS] [DE:HYPOTHETICAL 10.4 DRF3)] [SP:P32728] [DB:swissprot] [PN:conserved hypothetical protein [GN:ylxR] [OR:Bacillus subtilis] AC:Z18631] [GN:ORF3] [OR:Bacillus subtilis] Deperon.] [SP:P32728] [LE:2090] [RE:2365]
ORF Name A17503000983_4334383_f2_322 Description NO-HIT ORF Name A17503000983_4336536_c1_739 Description sp:[LN:YLXR_BACSU] [AC:P32728] [GN:YKD PROTEIN IN NUSA-INFB INTERGENIC R >pir:[LN:D36905] [AC:D36905:D69882:SylxR:hypothetical protein 1 (nusA 3'[DB:pir2] >gp:[GI:g580900] [LN:BSORF [DB:genpept-bct1] [DE:B.subtilis inf [DI:direct] >gp:[GI:e1185252:g263403]	NTID [580] LXR] [OF EGION (COMPANY) [APPROXIMATION (COMPANY) [APPROXI	AAID LengthLength AAID LengthLength AAID LengthLength LengthLength Score probability AAID LengthLength LengthLength LengthLength Score probability A352 315 104 211 3.3e-17 R:BACILLUS SUBTILIS] [DE:HYPOTHETICAL 10.4 DRF3)] [SP:P32728] [DB:swissprot] [PN:conserved hypothetical protein [GN:ylxR] [OR:Bacillus subtilis] AC:Z18631] [GN:ORF3] [OR:Bacillus subtilis] Deperon.] [SP:P32728] [LE:2090] [RE:2365] BSUB0009] [AC:Z99112:AL009126] [GN:ylxR]
ORF Name A17503000983_4334383_f2_322 Description NO-HIT ORF Name A17503000983_4336536_c1_739 Description sp:[LN:YLXR_BACSU] [AC:P32728] [GN:YKD PROTEIN IN NUSA-INFB INTERGENIC R >pir:[LN:D36905] [AC:D36905:D69882:SylxR:hypothetical protein 1 (nusA 3'[DB:pir2] >gp:[GI:g580900] [LN:BSORF [DB:genpept-bct1] [DE:B.subtilis inf [DI:direct] >gp:[GI:e1185252:g263403 [FN:unknown] [OR:Bacillus subtilis]	NTID [580] LXR] [OF EGION (COME STATE ST	AAID LengthLength AAID NT AA LengthLength LengthLength score probability AAID LengthLength AAID LengthLength LengthLength Below and a secore probability A352 315 104 211 3.3e-17 CR:BACILLUS SUBTILIS] [DE:HYPOTHETICAL 10.4 DRF3)] [SP:P32728] [DB:swissprot] [PN:conserved hypothetical protein CR:BACILLUS SUBTILIS] [OR:Bacillus subtilis] AC:Z18631] [GN:ORF3] [OR:Bacillus subtilis] AC:Z18631] [GN:ORF3] [OR:Bacillus subtilis] Deperon.] [SP:P32728] [LE:2090] [RE:2365] BSUB0009] [AC:Z99112:AL009126] [GN:ylxR] Dept-bct1] [DE:Bacillus subtilis complete
ORF Name A17503000983_4334383_f2_322 Description NO-HIT ORF Name A17503000983_4336536_c1_739 Description sp:[LN:YLXR_BACSU] [AC:P32728] [GN:YKD PROTEIN IN NUSA-INFB INTERGENIC R >pir:[LN:D36905] [AC:D36905:D69882:SylxR:hypothetical protein 1 (nusA 3'[DB:pir2] >gp:[GI:g580900] [LN:BSORF [DB:genpept-bct1] [DE:B.subtilis inf [DI:direct] >gp:[GI:e1185252:g263403 [FN:unknown] [OR:Bacillus subtilis]	NTID [580] [580] [580] [580] [580] [580] [580] [580] [680] [774] [78] [78] [78] [78] [78] [78] [78] [78] [78]	AAID LengthLength AAID NT AA LengthLength LengthLength score probability AAID LengthLength R:BACILLUS SUBTILIS] [DE:HYPOTHETICAL 10.4 DRF3)] [SP:P32728] [DB:swissprot] [PN:conserved hypothetical protein [GN:ylxR] [OR:Bacillus subtilis] AC:Z18631] [GN:ORF3] [OR:Bacillus subtilis] Deperon.] [SP:P32728] [LE:2090] [RE:2365] SUB0009] [AC:Z99112:AL009126] [GN:ylxR] Dept-bct1] [DE:Bacillus subtilis complete D7200.] [NT:alternate gene name: ymxB;
ORF Name A17503000983_4334383_f2_322 Description NO-HIT ORF Name A17503000983_4336536_c1_739 Description sp:[LN:YLXR_BACSU] [AC:P32728] [GN:YKD PROTEIN IN NUSA-INFB INTERGENIC R >pir:[LN:D36905] [AC:D36905:D69882:SylxR:hypothetical protein 1 (nusA 3'[DB:pir2] >gp:[GI:g580900] [LN:BSORF[DB:genpept-bct1] [DE:B.subtilis inf[DI:direct] >gp:[GI:e1185252:g263403] [FN:unknown] [OR:Bacillus subtilis] genome (section 9 of 21): from 15984	NTID [580] [580] [580] [580] [580] [580] [580] [580] [680] [774] [78] [78] [78] [78] [78] [78] [78] [78] [78]	AAID LengthLength AAID NT AA LengthLength LengthLength score probability AAID LengthLength R:BACILLUS SUBTILIS] [DE:HYPOTHETICAL 10.4 DRF3)] [SP:P32728] [DB:swissprot] [PN:conserved hypothetical protein [GN:ylxR] [OR:Bacillus subtilis] AC:Z18631] [GN:ORF3] [OR:Bacillus subtilis] Deperon.] [SP:P32728] [LE:2090] [RE:2365] SUB0009] [AC:Z99112:AL009126] [GN:ylxR] Dept-bct1] [DE:Bacillus subtilis complete D7200.] [NT:alternate gene name: ymxB;
ORF Name A17503000983_4334383_f2_322 Description NO-HIT ORF Name A17503000983_4336536_c1_739 Description sp:[LN:YLXR_BACSU] [AC:P32728] [GN:YKD PROTEIN IN NUSA-INFB INTERGENIC R >pir:[LN:D36905] [AC:D36905:D69882:SylxR:hypothetical protein 1 (nusA 3'[DB:pir2] >gp:[GI:g580900] [LN:BSORF[DB:genpept-bct1] [DE:B.subtilis inf[DI:direct] >gp:[GI:e1185252:g263403] [FN:unknown] [OR:Bacillus subtilis] genome (section 9 of 21): from 15984	NTID 580 LXR] [OF EGION (C 31992] region) 1T7A] [A B-nusA c 3] [LN:E [DB:geng 21to 180 [LE:134]	AAID LengthLength AAID NT AA LengthLength LengthLength score probability AAID LengthLength AAID LengthLength LengthLength ElengthLength AC: BACILLUS SUBTILIS] [DE: HYPOTHETICAL 10.4 DRF3)] [SP: P32728] [DB: swissprot] [PN: conserved hypothetical protein [GN: ylxR] [OR: Bacillus subtilis] AC: Z18631] [GN: ORF3] [OR: Bacillus subtilis] Depron.] [SP: P32728] [LE: 2090] [RE: 2365] BSUB0009] [AC: Z99112: AL009126] [GN: ylxR] Dept-bct1] [DE: Bacillus subtilis complete D7200.] [NT: alternate gene name: ymxB; A381] [RE: 134656] [DI: direct]
ORF Name A17503000983_4334383_f2_322 Description NO-HIT ORF Name A17503000983_4336536_c1_739 Description sp:[LN:YLXR_BACSU] [AC:P32728] [GN:YKD PROTEIN IN NUSA-INFB INTERGENIC R >pir:[LN:D36905] [AC:D36905:D69882:SylxR:hypothetical protein 1 (nusA 3'[DB:pir2] >gp:[GI:g580900] [LN:BSORF[DB:genpept-bct1] [DE:B.subtilis inf [DI:direct] >gp:[GI:e1185252:g263403 [FN:unknown] [OR:Bacillus subtilis] genome (section 9 of 21): from 15984 similar to hypothetical] [SP:P32728]	NTID [580] LXR] [OF EGION (C 31992] region) 1T7A] [A B-nusA c 3] [LN:E [DB:geng 21to 180 [LE:134] [LE:134]	AAID LengthLength AAID LengthLength LengthLength LengthLength LengthLength AAID LengthLength LengthLength Score probability A352 315 104 211 3.3e-17 R:BACILLUS SUBTILIS] [DE:HYPOTHETICAL 10.4 DRF3)] [SP:P32728] [DB:swissprot] [PN:conserved hypothetical protein [GN:ylxR] [OR:Bacillus subtilis] AC:Z18631] [GN:ORF3] [OR:Bacillus subtilis] Deperon.] [SP:P32728] [LE:2090] [RE:2365] SSUB0009] [AC:Z99112:AL009126] [GN:ylxR] Dept-bct1] [DE:Bacillus subtilis complete D7200.] [NT:alternate gene name: ymxB; A381] [RE:134656] [DI:direct]
ORF Name A17503000983_4334383_f2_322 Description NO-HIT ORF Name A17503000983_4336536_c1_739 Description sp:[LN:YLXR_BACSU] [AC:P32728] [GN:YKD PROTEIN IN NUSA-INFB INTERGENIC R >pir:[LN:D36905] [AC:D36905:D69882:SylxR:hypothetical protein 1 (nusA 3'[DB:pir2] >gp:[GI:g580900] [LN:BSORF[DB:genpept-bct1] [DE:B.subtilis inf [DI:direct] >gp:[GI:e1185252:g263403 [FN:unknown] [OR:Bacillus subtilis] genome (section 9 of 21): from 15984 similar to hypothetical] [SP:P32728]	NTID [580] LXR] [OF EGION (C 31992] region) 1T7A] [A B-nusA c 3] [LN:E [DB:genp 21to 180 [LE:134] [LE:134]	AAID LengthLength AAID NT AA LengthLength LengthLength score probability AAID LengthLength AAID LengthLength LengthLength ElengthLength AC: BACILLUS SUBTILIS] [DE: HYPOTHETICAL 10.4 DRF3)] [SP: P32728] [DB: swissprot] [PN: conserved hypothetical protein [GN: ylxR] [OR: Bacillus subtilis] AC: Z18631] [GN: ORF3] [OR: Bacillus subtilis] Depron.] [SP: P32728] [LE: 2090] [RE: 2365] BSUB0009] [AC: Z99112: AL009126] [GN: ylxR] Dept-bct1] [DE: Bacillus subtilis complete D7200.] [NT: alternate gene name: ymxB; A381] [RE: 134656] [DI: direct]

NO-HIT

ORF Name	NTID	AAID LengthLength score probability
A17503000983_4425068_£2_474	582	4354 2613 870 880 4.2e-88
subtilis] [DB:pir2] >gp:[GI:e1182850 [GN:yfhO] [FN:unknown] [OR:Bacillus statement of complete genome (section 5 of 21): fr [DI:direct] >gp:[GI:d1025397:g2804545]	g263318: subtilis rom 8028 [LN:I [DB:gen	al protein yfhO] [GN:yfhO] [OR:Bacillus 84] [LN:BSUB0005] [AC:Z99108:AL009126] 8] [DB:genpept-bct1] [DE:Bacillus subtilis 821 to1011250.] [LE:128691] [RE:131150] D85082] [AC:D85082] [PN:YfhO] [OR:Bacillus expept-bct1] [DE:Bacillus subtilis DNA, LE:21582] [RE:24041] [DI:direct]
ORF Name	NTID	AAID NT AA score probability
AI7503000983_4460063_f1_150 Description NO-HIT	583	4355 123 40
ORF Name AI7503000983_447326_c2_871 Description NO-HIT	<u>NTID</u>	AAID NT AA score probability 4356 156 51
ORF Name AI7503000983_4493778_c3_952 Description NO-HIT	<u>NTID</u>	AAID NT AA score probability 4357 48
[OR:Bacillus subtilis] [DB:pir2] >gp: [AC:Z99112:AL009126] [GN:yluC] [FN:ur	:[GI:e1] nknown] e (secti	[OR:Bacillus subtilis] [DB:genpept-bct1] ion 9 of 21): from 1598421to 1807200.]
ORF Name A17503000983_4687825_c3_955 Description NO-HIT	NTID 587	AAID NT AA score probability 4359 165 54
ORF Name A17503000983_4719011_c2_841 Description NO-HIT	<u>NTID</u> 	AAID NT AA score probability 4360 174 57
ORF Name AI7503000983_4740932_c1_825 Description NO-HIT	<u>NTID</u> 589	AAID NT AA score probability 4361 126 41

 $\underline{\mathtt{NT}}$

<u> AA</u>

ORF Name	NTID	AAID	<u>NT</u> LengthI	<u>AA</u> Length	score	probability	
AI7503000983_4798453_c1_790	590	4362	1026	341	242	2.7e-27	
Description pir:[LN:H69873] [AC:H69873] [PN:c [OR:Bacillus subtilis] [DB:pir2] > [PN:YlbC protein] [GN:ylbC] [OR:Ba subtilis genomic DNA 23.9kB fragme >gp:[GI:e1185086:g2633867] [LN:BSU [OR:Bacillus subtilis] [DB:genpept 8 of 21): from 1394791to 1603020.] subtilis] [LE:170993] [RE:172033]	egp:[GI:ei acillus su ent.] [LE JB0008] [A :-bct1] [I [NT:sim	334771: ubtilis :11510] AC:Z991 DE:Baci ilar to	g2339999] [DB:ge [RE:125 11:AL009 llus sub	9] [LN enpept 550] [9126] otilis	I:BS168 -bct1] [DI:din [GN:y] comp]	B23KB] [AC:Z98682] [DE:Bacillus rect] bC] [FN:unknown] Lete genome (sectio	n
ORF Name	NTID	AAID	<u>NT</u> LengthL	<u>AA</u> Length	score	probability	_
AI7503000983_4884675_c3_1057	591	4363	1212	403	1268	3.2e-129	
sp:[LN:TRPB_LACLA] [AC:Q01998] [GN [SR:,SUBSPLACTIS:STREPTOCOCCUS LAC CHAIN,] [SP:Q01998] [DB:swissprot] synthase, beta chain] [GN:trpB] [beta chain homology] [OR:Lactococco pgp:[GI:g149521] [LN:LACTRPOP] [AC [GN:trpB] [OR:Lactococcus lactis] lactis) DNA] [DB:genpept-bct1] [EC trpC, trpB trpA genes, completecds	TIS] [EC pir:[LN pir:[:4.2.1. N:S3512 ophan s s subsp [PN:tr ococcus O] [DE:	20] [DE: 9] [AC:S ynthase . lactis yptophar lactis L. lacti	:TRYPT 535129 beta s] [EC n synt (stra is trp	OPHAN Property Property	N:tryptophan ctryptophan synthas 1.20] [DB:pir2] peta subunit] 1403, sub_species pG, trpD, trpF,	е
ORF Name	NTID	AAID	<u>NT</u> LengthL	<u>AA</u> .ength	score	probability	
A17503000983_4890802_f2_275	592	4364	1152		100	1.2e-81	
Description sp:[LN:TYRA_BACSU] [AC:P20692] [GN [DE:PREPHENATE DEHYDROGENASE, (PDH						:1.3.1.12]	
ORF Name A17503000983 4891577 c2 879	NTID	AAID	NT LengthL				
Description pir: [LN:G69657] [AC:G69657] [PN:t [GN:miaA] [CL:delta(2)-isopenteny [DB:pir2] >gp:[GI:e1183392:g263411 isopentenylpyrophosphate transfera [DB:genpept-bct1] [DE:Bacillus sub 1781201to 2014980.] [LE:84430] [RE	rlpyrophos 7] [LN:BS se] [GN:n stilis con	sphate SUB0010 miaA] [mplete	pyrophos transfer [AC:Z9 OR:Bacil genome (case] 99113: Llus s	trans [OR:Ba AL0091 ubtili	cillus subtilis] .26] [PN:tRNA .s]	
ORF Name A17503000983_4942202_c3_1058 Description	<u>NTID</u>	<u>AAID</u>	NT LengthL	AA ength	score	probability	_

NO-HIT

ODE Name	NMTD	7 7 T D	NT	AA			-3176
ORF Name	NTID	AAID	Length:	Length	score	probar	oility
A17503000983_4964686_t3_520	595	4367	204	67	195	1.6e-15	
Description							
pir: [LN:C70057] [AC:C70057] [PN:4-o [CL:4-oxalocrotonate tautomerase] [O >gp: [GI:e267624:g1565237] [LN:BSTHRZ Pseudomonas putida] [GN:ywhB] [OR:Ba thrZ downstream chromosomal region.] >gp: [GI:e1186254:g2636290] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-b 20 of 21): from 3798401to 4010550.] [LE:54138] [RE:54326] [DI:direct] ORF Name AI7503000983_5109378_f3_546 Description pir: [LN:A69892] [AC:A69892] [PN:con	R:Bacill [AC:Z8 cillus s [LE:238 020] [AC ct1] [DE [NT:simi	lus suk 30360] subtili 38] [RE C:Z9912 E:Bacil ilar to AAID 4368	PN:Un S [DB S:2576] C:2576] C:4-oxa NT Length	[DB:pknown,:genpe [DI:c 9126] btilis locrot AA Length 204	highl highl pt-bct complen [GN:yw compl conate score 469 n ynes	y simila [DE:I nent] hB] [FN ete gend tautomen probak	ar to B.subtilis :unknown] ome (section rase]
[CL:Escherichia coli ygiH protein] [0 > gp:[GI:e249655:g1405459] [LN:BC170D] subtilis] [DB:genpept-bct1] [DE:B.suf [NT:similar to hypothetical protein if > gp:[GI:e1183465:g2634190] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-bct0] of 21): from 1781201to 2014980.] [RE:150556] [DI:complement]	EGR] [AC btilis I MG247 fr 010] [AC ct1] [DE	C:Z7323 DNA (26 com] [L C:Z9911 E:Bacil	4] [PN .2 kb .E:1359 .3:AL00 .lus su	:YneS] fragme 6] [RE 9126] btilis	[GN:ynt; 17 ::14177 [GN:yn	0 degree] [DI:co es] [FN: ete geno	e region).] omplement] unknown] ome (section
ORF Name	NTID	AAID	<u>NT</u> Length1	<u>AA</u> Length	score	probab	oility
AI7503000983_5109625_c3_1011	597	4369		123	258	3.4e-22	
Description		L			الــــــــا		
pir:[LN:D70039] [AC:D70039] [PN:two [GN:yvfU] [CL:regulatory protein consubtilis] [DB:pir2] >gp:[GI:e1186094] [GN:yvfU] [FN:unknown] [OR:Bacillus accomplete genome (section 18 of 21): two-component response regulator [Yv: >gp:[GI:e313075:g1945721] [LN:BSZ9404] [OR:Bacillus subtilis] [DB:genpept-backet] [NT:probable two component regulator]	mA:respo :g263591 subtilis from 339 fT]] [LE 43] [AC: ct1] [DE	onse re .9] [LN :] [DB: :9551to :95389 :294043	gulato: :BSUB0 genpep 36090 [RE::] [PN:]	r homo 018] [t-bct1 60.] [95991] hypoth genomi	logy] AC:Z99] [DE: NT:sim [DI:c etical c DNA	[OR:Baci 121:AL00 Bacillus ilar to omplemer proteir fragment	illus 09126] s subtilis nt] n] [GN:yvfU] c (88 kb).]
ORF Name	NTID	AAID	<u>NT</u> LengthI	<u>AA</u> Length	score	probab	ility
A17503000983_5120635_c2_903 Description	598	-		79	76	0.026	
sp:[LN:F801_SCHMA] [AC:P16463] [OR:S0 SPECIFIC 800 PROTEIN (FS800)] [SP:P16 [AC:J03999] [PN:female-specific 800 p [SR:Schistosoma mansoni (strain Puero [DE:Schistosoma mansoni female-specific [NT:putative] [LE:4] [RE:720] [DI:din	5463] [D protein] to Rican fic 800	B:swis [GN:f CDNA	sprot] s800] to mR1	>gp:[0 [OR:Sc] NA] [D]	GI:g16 histos B:genp	0990] [I oma mans ept-invl	LN:SCMFS800] soni] .]

A17503000983_5195328_c1_749 Description gp:[GI:g1842438] [LN:BSU87792] [AC:U87 [DB:genpept-bct1] [DE:Bacillus subtili	NTID AAID LengthLength score probability 599 4371 393 130 142 2.3e-09 7792] [PN:unknown] [OR:Bacillus subtilis] is tRNA-Ala, phosphatidylglycerophosphate complete cds, and RecA (recA) gene, partial cds]
ORF Name AI7503000983_5198557_c1_734 Description gp:[GI:d1032955:g3358087] [LN:AB004319	NTID AAID NT AA Score probability 600 4372 801 266 847 1.3e-84 9] [AC:AB004319] [PN:undecaprenyl diphosphate uteus] [SR:Micrococcus luteus (strain:B-P 26) I eus DNA for undecaprenyl diphosphate] NA]
Description pir:[LN:F69880] [AC:F69880] [PN:conse [CL:conserved hypothetical protein MG4 >gp:[GI:e1185196:g2633977] [LN:BSUB000 [OR:Bacillus subtilis] [DB:genpept-bct	NTID AAID NT AA Score probability LengthLength Score probability 4373 867 288 839 9.2e-84 erved hypothetical protein ylqF] [GN:ylqF] 442] [OR:Bacillus subtilis] [DB:pir2] 09] [AC:Z99112:AL009126] [GN:ylqF] [FN:unknown] t1] [DE:Bacillus subtilis complete genome (sect T:similar to hypothetical proteins] [LE:77511]	
A17503000983_5355012_c1_770 [6] Description sp:[LN:GLNA_STAAU] [AC:Q59812] [GN:GLN [DE:GLUTAMINE SYNTHETASE, (GLUTAMATE >gp:[GI:e214721:g1134886] [LN:SAGLNAR]	NTID AAID LengthLength score probability LengthLength Score Probability LengthLength Length Length Length Length LengthLength Length Length Length Length LengthLength Length Length Length Length LengthLength Length Length Length Length Length Length Length Length Length Length Length Length Length	A]
	NTID AAID NT AA score probability 603 4375 153 50	

Description NO-HIT

ORF Name probability A17503000983 6258588 cl 802 604 1489 1305 Description sp:[LN:ALST BACSU] [AC:Q45068] [GN:ALST] [OR:BACILLUS SUBTILIS] [DE:AMINO ACID CARRIER PROTEIN ALST] [SP:Q45068] [DB:swissprot] >pir:[LN:A69585] [AC:A69585] [PN:amino acid carrier protein alsT] [GN:alsT] [CL:sodium-dependent D-alanine/qlycine transport protein] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e249660:g1405464] [LN:BC170DEGR] [AC:Z73234] [PN:AlsT] [GN:alsT] [FN:aminoacid carrier protein] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:B.subtilis DNA (26.2 kb fragment; 170 degree region).] [NT:similar to sodium/proton dependent alanine carrier] [SP:Q45068] [LE:20601] [RE:21998] [DI:direct] >qp:[GI:e1183470:q2634195] [LN:BSUB0010] [AC:Z99113:AL009126] [PN:amino acid carrier protein] [GN:alsT] [OR:Bacillus subtilis] [DB:qenpept-bct1] [DE:Bacillus subtilis complete genome (section 10 of 21): from 1781201to 2014980.] [SP:Q45068] [LE:156980] [RE:158377] [DI:direct] AΑ ORF Name NTID AAID <u>score</u> probability LengthLength AI7503000983 6416566 cl 800 605 2031 3180 Description sp:[LN:PARE_STAAU] [AC:P50072] [GN:PARE:GRLB] [OR:STAPHYLOCOCCUS AUREUS] [EC:5.99.1.-] [DE:TOPOISOMERASE IV SUBUNIT B,] [SP:P50072] [DB:swissprot] >pir:[LN:S54426] [AC:S54426] [PN:DNA topoisomerase (ATP-hydrolyzing), chain B] [CL:DNA topoisomerase (ATP-hydrolyzing) chain B] [OR:Staphylococcus aureus] [EC:5.99.1.3] [DB:pir2] >gp:[GI:d1011746:g1777320] [LN:D67075] [AC:D67075] [PN:DNA topoisomerase IV GrlB subunit] [GN:grlB] [OR:Staphylococcus aureus] [SR:Staphylococcus aureus (isolate:RN4220) DNA] [DB:genpept-bct1] [DE:Staphylococcus aureus DNA for DNA topoisomerase IV GrlB subunit, DNA topoisomerase IV GrlA subunit, complete cds.] [LE:385] [RE:2376] [DI:direct] >gp:[GI:g561879] [LN:STAGYRASL] [AC:L25288] [PN:gyrase-like protein beta subunit] [GN:grlB] [OR:Staphylococcus aureus] [SR:Staphylococcus aureus (tissue library: FDA 574) DNA] [DB:genpept-bct1] [DE:Staphylococcus aureus gyrase-like protein alpha and beta subunit (grlA and grlB) genes, complete cds.] [LE:41] [RE:2032] [DI:direct] >gp:[GI:e306312:g2302281] [LN:A48501] [AC:A48501] [OR:Staphylococcus aureus] [DB:genpept-pat] [DE:Sequence 3 from Patent WO9603516.] [NT:unnamed protein product] [LE:1] [RE:1992] [DI:direct]

 ORF Name
 NTID
 AAID
 NT AA LengthLength
 Score
 probability

 A17503000983_6525_f3_577
 606
 4378
 1035
 344
 729
 4.2e-72

Description

sp:[LN:LYSP_ECOLI] [AC:P25737] [GN:LYSP:CADR] [OR:ESCHERICHIA COLI] [DE:LYSINE-SPECIFIC
PERMEASE] [SP:P25737] [DB:swissprot]

ORF Name	NTID	AAID	NT	<u>AA</u>	zaoro	probability
A17503000983_6641963_c3_1009			Length Le			
	607	4379	1485 4	94	1302	8.0e-133
Description sp:[LN:YWNE_BACSU] [AC:P71040] [GN:YKD PROTEIN IN SPOIIQ-MTA INTERGENIC [AC:G70063] [PN:cardiolipin synthas cardiolipin synthetase] [OR:Bacillus [LN:BSUB0019] [AC:Z99122:AL009126] [DB:genpept-bct1] [DE:Bacillus subti 3597091to 3809700.] [NT:similar to c [RE:166076] [DI:direct] >gp:[GI:e269 [GN:ywnE] [OR:Bacillus subtilis] [DE downstream DNA.] [NT:Product similar [LE:5155] [RE:6603] [DI:complement] [AC:Z99122:AL009126] [GN:ywnE] [FN:ume] [DE:Bacillus subtilis complete genome [NT:similar to cardiolipin synthase]	REGION] se homolo s subtil [GN:ywnE lis comp eardioli 9549:g15 3:genpep t to Esc >gp:[GI unknown] ne (sect	[SP:Pog ywn is] [D] [FN:plete pin sy 92701] t-bct1 herich [OR:B ion 19	71040] [IE] [GN:your B:pir2] : unknown] genome (such ase] [LN:BSUI] [DE:B.suia coli coli coli coli coli coli coli coli	DB:swi wnE] >gp:[G [OR:E section [SP:P7 EROP] subtil cardion 6184] subtil from	Esspro [CL:E [CL:E EI:ell Bacill on 19 71040] [AC:Y Lis ur blipin [LN:E Lis] [printer [LN:G70063] cacillus probable 84565:g2636184] us subtilis] of 21): from [LE:164628] 08559] [PN:Unknown] case operon and [SP:P71040] SUB0019] DB:genpept] 91to 3809700.]
ORF Name	NTID	AAID	<u>NT</u> LengthLe	AA ength	score	probability
A17503000983_6664127_c2_940	608	4380	438 14	45		
Description						
NO-HIT						
ORF Name	NTID	AAID	NT	AA s	score	probability
AI7503000983_6678140_c3_972	609	4381	LengthLe			3.8e-28
Description				1		
pir:[LN:S52267] [AC:S52267] [PN:DNA III alpha chain polC] [OR:Staphyloco [LN:SADNAPOL3] [AC:Z48003:L39156] [F [DB:genpept-bct1] [DE:S.aureus gene [RE:4281] [DI:direct]	occus au N:DNA po	reus] olymer	[DB:pir2] ase III]	>gp: OR:S	[GI:g staphy	642270] lococcus aureus]
ORF Name	NTID	AAID	NT LengthLe	AA ength	core	probability
A17503000983_6688126_c1_751	610	4382	1074 3			7.2e-171
Description	_					·
sp:[LN:RECA_STAAU] [AC:Q02350] [GN:R [SP:Q02350] [DB:swissprot] >gp:[GI:g	463285]	[LN:S'	TARECAA]	[AC:L	25893] [GN:recA]
[FN:genetic recombination] [OR:Staph [DB:genpept-bct1] [DE:Staphylococcus [LE:16] [RE:1059] [DI:direct]						
[DI.GIECC]	and the second s			And the second Suprement		The state of the s
ORF Name	NTID	AAID	NT LengthLe	AA ength	core	probability
MTTENANNORS CTAVAT FO FOR						
A17503000983_673437_f3_529	611	4383	147 48			

NO-HIT

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000983_6818827_c3_1072	612	4384	1608	535	2088	4.1e-216
Description	·	L	ــــاك		<u> </u>	
pir:[LN:E69861] [AC:E69861] [PN:ABC [GN:ykpA] [CL:ATP-binding cassette pgp:[GI:e1185033:g2633814] [LN:BSUBC [OR:Bacillus subtilis] [DB:genpept-k B of 21): from 1394791to 1603020.] [protein)] [LE:116988] [RE:118610] [I [AC:AF012285:AF012284:U51911] [PN:Yk [DB:genpept-bct2] [DE:Bacillus subtications of the support of the	homolog 0008] [A oct1] [D [NT:simi DI:direc apA] [GN Llis mob	y] [OR C:Z991 E:Baci lar to t] >gp :ykpA] A-nprE	:Bacill 11:AL00 llus su ABC tr :[GI:g3 [OR:Ba	us sub 09126] obtilis canspor 3282128 ocillus region.	tilis [GN:y comp ter (] [LN subt] [NT] [DB:pir2] kpA] [FN:unknown] lete genome (section ATP-binding :AF012285] ilis]
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
17503000983_6837812_c1_735	613	4385	786	261	611	1.3e-59
Description						
sp:[LN:CDSA_BACSU] [AC:O31752] [GN:C [DE:SYNTHASE)] [SP:O31752] [DB:swiss [PN:phosphatidate cytidylyltransfera [DB:pir2] >gp:[GI:e1185245:g2634026] [PN:phosphatidate cytidylyltransfera [OR:Bacillus subtilis] [DB:genpept-k genome (section 9 of 21): from 15984 [DI:direct]	sprot] > se cdsA [LN:BS se] [GN oct1] [E	pir:[L]] [GN: UB0009 :cdsA] C:2.7.	N:G6959 cdsA]] [AC:Z [FN:ph 7.41] [7] [AC [OR:Ba 399112: losphol DE:Bac	:G695; cillu: AL009; ipid l illus	97] s subtilis] 126] biosynthesis] subtilis complete
DRF Name	NTID	AAID	NT Length	<u>AA</u> Length	score	probability
17503000983_6929652_c1_822	614	4386		247	683	3.1e-67
Description pir: [LN:F69866] [AC:F69866] [PN:tet [GN:ykuQ] [OR:Bacillus subtilis] [I [AC:AJ222587] [PN:YkuQ protein] [GN: [DE:Bacillus subtilis 29kB DNA fragm to acetyltransferases] [LE:23332] [R [LN:BSUB0008] [AC:Z99111:AL009126] [I [DB:genpept-bct1] [DE:Bacillus subtilis 294791to 1603020.] [NT:similar to t [RE:94298] [DI:direct]	OB:pir2] ykuQ] [(nent from EE:24042] GN:ykuQ Lis com	>gp:[OR:Bac m ykwC] [DI:] [FN: plete	GI:el18 illus s gene t direct] unknown genome	1922:g ubtili o csel >gp:[] [OR:	263223 s] [DI 5 gene GI:ell Bacill on 8 d	B8] [LN:BS16829KB] B:genpept-bct1] e.] [NT:homologous 185008:g2633789] lus subtilis] of 21): from
DRF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
17503000983_6929677_c1_766	615	4387	258	85	187	1.1e-14
Description						
oir:[LN:B69884] [AC:B69884] [PN:hos [CL:host factor I] [OR:Bacillus subt						

[LN:BSUB0010] [AC:Z99113:AL009126] [GN:ymaH] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 10 of 21): from 1781201to 2014980.] [NT:similar to host factor-1 protein] [LE:85414] [RE:85635] [DI:direct]

ORF Name probability

AI7503000983_6933390_c1_759

616

4388

183

1552

1.9e-37

Description

sp:[LN:GLPP_BACSU] [AC:P30300] [GN:GLPP] [OR:BACILLUS SUBTILIS] [DE:GLYCEROL UPTAKE OPERON ANTITERMINATOR REGULATORY PROTEIN] [SP:P30300] [DB:swissprot] >pir:[LN:B47700] [AC:B47700:D69634] [PN:glycerol metabolism regulatory protein GlpP] [GN:glpP] [OR:Bacillus subtilis] [DB:pir2] [MP:75 (degrees)] >gp:[GI:g142996] [LN:BACGLPPFK] [AC:M99611] [PN:regulatory protein] [GN:glpP] [FN:putative antiterminator] [OR:Bacillus subtilis] [SR:Bacillus subtilis DNA] [DB:genpept-bct1] [DE:Bacillus subtilis antiterminator regulatory protein (glpP), glycerol uptake facilitator (glpF) genes, complete cds, glycerolkinase (glpK) gene, 5' end.] [LE:328] [RE:906] [DI:direct] >gp:[GI:e1182916:g2633250] [LN:BSUB0005] [AC:Z99108:AL009126] [PN:transcription antiterminator] [GN:glpP] [FN:control of mRNA stability of glpD] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 5 of 21): from 802821 tol011250.] [SP:P30300] [LE:198429] [RE:199007] [DI:direct] >gp:[GI:e1182928:q2633262] [LN:BSUB0006] [AC:Z99109:AL009126] [PN:transcription antiterminator] [GN:glpP] [FN:control of mRNA stability of glpD] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 6 of 21): from 999501 to1209940.] [SP:P30300] [LE:1749] [RE:2327] [DI:direct] >gp:[GI:e324939:g2226135] [LN:BSY14079] [AC:Y14079] [PN:regulatory protein] [GN:glpP] [FN:putative antiterminator] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis chromosomal DNA, region 75 degrees: glpPFKDoperon and downstream.] [NT:see EMBL M99611 and Swiss Prot P30300.] [SP:P30300] [LE:1397] [RE:1975] [DI:direct]

NT AA ORF Name NTID AAID score probability LengthLength AI7503000983_7064077_c1_740 4389 2187 728 1.3e-260 617

Description

sp:[LN:IF2 BACSU] [AC:P17889:031757] [GN:INFB] [OR:BACILLUS SUBTILIS] [DE:TRANSLATION INITIATION FACTOR IF-2] [SP:P17889:O31757] [DB:swissprot] >pir:[LN:A35269] [AC:A35269:B35269:S31994:G69644] [PN:translation initiation factor IF-2] [GN:infB] [CL:translation initiation factor IF-2:translation elongation factor Tu homology] [OR:Bacillus subtilis] [DB:pir1] >gp:[GI:g143359] [LN:BACPSIF2A] [AC:M34836] [OR:Bacillus subtilis] [SR:B.subtilis (strain RS410) DNA, clones lambda-JET[1,2],pUK, an] [DB:genpept-bct1] [DE:B.subtilis protein synthesis initiation factor 2 (infB) gene, complete cds.] [NT:protein synthesis initiation factor 2 (infB)] [LE:381] [RE:2531] [DI:direct] >gp:[GI:g49319] [LN:BSORF1T7A] [AC:Z18631] [GN:IF2] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:B.subtilis infB-nusA operon.] [SP:P17889] [LE:2689] [RE:4839] [DI:direct]

NT AΑ ORF Name NTID AAID <u>score</u> probability LengthLength AI7503000983_7203176_c1_810 618 4390 801 266 426 5.4e-40

Description

sp:[LN:TRPA METJA] [AC:Q60180] [GN:TRPA:MJ1038] [OR:METHANOCOCCUS JANNASCHII] [EC:4.2.1.20] [DE:TRYPTOPHAN SYNTHASE ALPHA CHAIN,] [SP:Q60180] [DB:swissprot] >pir:[LN:E64429] [AC:E64429] [PN:tryptophan synthase, alpha chain] [CL:tryptophan synthase alpha chain:tryptophan synthase alpha chain homology] [OR:Methanococcus jannaschii] [EC:4.2.1.20] [DB:pir2] [MP:FOR969735-970589] >gp:[GI:g1591691] [LN:U67546] [AC:U67546:L77117] [PN:tryptophan synthase alpha subunit (trpA)] [GN:MJ1038] [OR:Methanococcus jannaschii] [DB:qenpept-bct2] [DE:Methanococcus jannaschii section 88 of 150 of the complete genome.] [NT:similar to GB:M65060 SP:P26920 PID:149750] [LE:5837] [RE:6691] [DI:direct]

ORF Name	NTID	AAID	NT AA LengthLength score probability	
AI7503000983_7242250_c2_924	619	4391		
Description	 			
gp:[GI:g5002553] [LN:AF074603] [AC:	\F074603`	1 [DN].	NonEl [GN:nonEl [OR:Streptomyces	
griseus subsp. griseus] [DB:genpept-				
			ace.] [LE:12384] [RE:13088] [DI:direct	t.1
ORF Name	NTID	AAID	NT AA LengthLength score probability	
A17503000983_801552_t2_228	620	4392	381 126 74 0.011	
Description				
gp:[GI:g2454643] [LN:AF020905] [AC:A	F020905	l [bw.	E5] [GN·E5] [OR:common chimpanzee	
papillomavirus 1] [DB:genpept-vrl]				
genome.] [LE:3901] [RE:4185] [DI:dir			mpanio presidenti di comprese	
ORF Name	NTID	AAID	NT AA score probability	
AI7503000983_835252_f1_13	621	4393		
Description		L		
NO-HIT				
	The section with the section of the			
ORF Name	NTID	AAID	NT AA score probability	
AI7503000983_837550_c2_914	622	4394	246 81	
Description				
NO-HIT				
				المرابعة
ORF Name	NTID	AAID	NT AA LengthLength score probability	
AI7503000983_869052_c2_876	623	4395	1677 558 1700 5.3e-175	
Description				
sp:[LN:GLPD_BACSU] [AC:P18158] [GN:G	LPD] [OF	R:BACI	LLUS SUBTILIS] [EC:1.1.99.5]	
[DE:AEROBIC GLYCEROL-3-PHOSPHATE DEH	YDROGEN	ASE,]	[SP:P18158] [DB:swissprot]	
>pir:[LN:C45868] [AC:C45868:A69634:S				
<pre>glpD] [GN:glpD] [OR:Bacillus subtil</pre>			_	
[LN:BACGLPKD] [AC:M34393] [OR:Bacill				
[DB:genpept-bct1] [DE:B.subtilis gly			5 2	
	_		omplete cds.] [NT:glycerol-3-phosphat	:e
dehydrogenase (glpD) (EC] [LE:2329]				
[LN:BSUB0005] [AC:Z99108:AL009126] [FN:glycerol utilization] [OR:Bacill				
[DE:Bacillus subtilis complete genom				
[SP:P18158] [LE:201660] [RE:203327]				
[LN:BSUB0006] [AC:Z99109:AL009126] [<u> </u>	
[FN:glycerol utilization] [OR:Bacill				
[DE:Bacillus subtilis complete genom				
[SP:P18158] [LE:4980] [RE:6647] [DI:				
[AC:Y14079] [PN:glycerol-3-phosphate	dehydro	ogenas	e] [GN:glpD] [OR:Bacillus subtilis]	
[DB:genpept-bct1] [DE:Bacillus subti			· •	
	ee EMBL	M3439	3 and Swiss Prot P18158.] [SP:P18158]	
[LE:4628] [RE:6295] [DI:direct]				

ORF Name	NTID AA	ID LengthLength score probability
A17503000983_891700_c2_865	624 43	96 867 288 599 2.5e-58
beta chain] [OR:Halobacterium halobi	um] [EC:1.2	
[DB:genpept-bct1] [EC:1.2.7.1] [DE:H	.halobium q	reductase] [OR:Halobacterium halobium] gene for pyruvate:ferredoxin nthase] [LE:2057] [RE:2995] [DI:direct]
ORF Name	NTID AA	ID LengthLength score probability
AI7503000983_892186_£1_171	625 43	97 [129] 42
Description NO-HIT		
ORF Name	NTID AA	ID <u>NT AA</u> LengthLength score probability
A17503000983_893826_t2_384 Description	626 43:	98 123 40 49 0.031
56 of 134 of the complete genome.] [:pir2] >gp hypothetic enpept-bct2	:[GI:g2313764] [LN:AE000578]
[LE:6414] [RE:6707] [DI:direct]	none with a control of management and a control of	
ORF Name	NTID AA	ID NT AA probability
ORF Name AI7503000983_9767263_c2_866 -	NTID AA.	LengthLength probability
ORF Name A17503000983_9767263_c2_866 Description pir: [LN:A69922] [AC:A69922] [PN:pha [GN:yoqZ] [CL:phage-related replica >gp: [GI:e1185518:g2634439] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-b 11 of 21): from 2000171to 2207900.] [RE:189856] [DI:complement] >gp: [GI: [OR:Bacteriophage SPBc2] [DB:genpept	ge-related tion protein [AC:25 ct1] [DE:Ba [NT:similar g3025599] [DE:F	LengthLength score probability 99 639 212 280 1.6e-24 replication protein homolog yoqZ]
ORF Name A17503000983_9767263_c2_866 Description pir: [LN:A69922] [AC:A69922] [PN:pha [GN:yoqZ] [CL:phage-related replica >gp: [GI:e1185518:g2634439] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-b 11 of 21): from 2000171to 2207900.] [RE:189856] [DI:complement] >gp: [GI: [OR:Bacteriophage SPBc2] [DB:genpept	ge-related tion protein [AC:25 ct1] [DE:Ba [NT:similar g3025599] [DE:F	LengthLength score probability 19 639 212 280 1.6e-24 replication protein homolog yoqZ] in] [OR:Bacillus subtilis] [DB:pir2] 1914:AL009126] [GN:yoqZ] [FN:unknown] 1018:Acillus subtilis complete genome (section or to phage-related protein] [LE:189032] [LN:AF020713] [AC:AF020713] [GN:yoqZ] 1028:Bacteriophage SPBc2 complete genome.] 1029:Inct [LE:95664] [RE:96488] [DI:direct]
ORF Name A17503000983_9767263_c2_866 Description pir: [LN:A69922] [AC:A69922] [PN:pha [GN:yoqZ] [CL:phage-related replica >gp: [GI:e1185518:g2634439] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-b 11 of 21): from 2000171to 2207900.] [RE:189856] [DI:complement] >gp: [GI: [OR:Bacteriophage SPBc2] [DB:genpept [NT:similar to bacteriophage SPP1 OR ORF Name A17503000983_978450_c1_804	ge-related tion protein [AC:ZS ct1] [DE:Ba [NT:similar g3025599] -phg] [DE:F37.1 produced]	LengthLength score probability 10 LengthLength score probability 11 LengthLength score probability 12 LengthLength score probability 13 LengthLength score probability 14 LengthLength score probability 15 LengthLength score probability 16 LengthLength score probability
ORF Name A17503000983_9767263_c2_866 Description pir: [LN:A69922] [AC:A69922] [PN:pha [GN:yoqZ] [CL:phage-related replica >gp:[GI:e1185518:g2634439] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-b 11 of 21): from 2000171to 2207900.] [RE:189856] [DI:complement] >gp:[GI: [OR:Bacteriophage SPBc2] [DB:genpept [NT:similar to bacteriophage SPP1 OR	ge-related tion protein [DE:Base [NT:similar g3025599] [DE:F37.1 produant [DE:Base [NT:D] AA: [Carter [DE:Base	LengthLength score probability 10 LengthLength score probability 11 LengthLength score probability 12 Polication protein homolog yoqZ] 13 [OR:Bacillus subtilis] [DB:pir2] 14 [OR:Bacillus subtilis] [FN:unknown] 15 [OR:Bacillus subtilis] [SN:yoqZ] 16 [CH:AF020713] [AC:AF020713] [GN:yoqZ] 17 [CH:Pose Probability 18 [CH:Pose Probability 19 [OR:Bacillus subtilis] 10 [OR:Bacillus subtilis] 10 [OR:Bacillus subtilis]
ORF Name A17503000983_9767263_c2_866 Description pir: [LN:A69922] [AC:A69922] [PN:pha [GN:yoqZ] [CL:phage-related replica >gp: [GI:e1185518:g2634439] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-b 11 of 21): from 2000171to 2207900.] [RE:189856] [DI:complement] >gp: [GI: [OR:Bacteriophage SPBc2] [DB:genpept [NT:similar to bacteriophage SPP1 OR ORF Name A17503000983_978450_c1_804 Description gp: [GI:d1025380:g2804528] [LN:D85082 [SR:Bacillus subtilis DNA] [DB:genpept	ge-related tion protein [AC:Z9 ct1] [DE:Ba	LengthLength 10 LengthLength 11 LengthLength 12 LengthLength 13 LengthLength 14 LengthLength 15 LengthLength 16 LengthLength 17 LengthLength 18 LengthLength 19 LengthLength 10 LengthLength 10 LengthLength 10 LengthLength 10 LengthLength 10 LengthLength 10 LengthLength 11 LengthLength 12 LengthLength 13 LengthLength 14 LengthLength 15 LengthLength 16 LengthLength 17 LengthLength 18 LengthLength 19 LengthLength 19 LengthLength 10 LengthLength 11 LengthLength 12 LengthLength 13 LengthLength 14 LengthLength 15 LengthLength 16 LengthLength 17 LengthLength 18 LengthLength 19 LengthLength 19 LengthLength 19 LengthLength 10 LengthLength 10 LengthLength 11 LengthLength 12 LengthLength 13 LengthLength 14 LengthLength 15 LengthLength 16 LengthLength 17 LengthLength 18 LengthLength 19 LengthLength 19 LengthLength 10 LengthLength 10 LengthLength 11 LengthLength 12 LengthLength 13 LengthLength 14 LengthLength 15 LengthLength 16 LengthLength 17 LengthLength 18 LengthLength 18 LengthLength 19 LengthLength 19 LengthLength 20 Length
ORF Name A17503000983_9767263_c2_866 Description pir: [LN:A69922] [AC:A69922] [PN:pha [GN:yoqZ] [CL:phage-related replica >gp:[GI:e1185518:g2634439] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-b 11 of 21): from 2000171to 2207900.] [RE:189856] [DI:complement] >gp:[GI:[OR:Bacteriophage SPBc2] [DB:genpept [NT:similar to bacteriophage SPP1 OR ORF Name A17503000983_978450_c1_804 Description gp:[GI:d1025380:g2804528] [LN:D85082 [SR:Bacillus subtilis DNA] [DB:genpe sequence, 79 to 81 degree region.] [ge-related tion protes of the	LengthLength Length LengthLength LengthLength LengthLength LengthLength Length LengthLength Length

ORF Name	NTID	AAID	NT Lengthl	<u>AA</u> Length	score	probability
A17503000983_9804202_c1_732	630	4402	627	208	704	1.9e-69
Description			_ L		· L	
pir:[LN:B69727] [AC:B69727] [PN:tra [CL:translation elongation factor EF >gp:[GI:e1185241:g2634022] [LN:BSUB0 [GN:tsf] [OR:Bacillus subtilis] [DB: (section 9 of 21): from 1598421to 18 [DI:direct]	'-Ts] [O: 009] [A genpept	R:Baci C:Z991 -bct1]	llus sul 12:AL00 DE:Ba	btilis 9126] cillus	DB: [PN:e] subti	pir2] Longation factor Ts] Llis complete genome
ORF Name	NTID	AAID	<u>NT</u> LengthI	<u>AA</u> Length	score	probability
AI7503000983_9807807_c1_748	631	4403	873	290	522	3.6e-50
Description gp:[GI:g1842437] [LN:BSU87792] [AC:U [DB:genpept-bct1] [DE:Bacillus subti synthase(pgsA) and CinA (cinA) genes [NT:hypothetical 17.9 kDa protein; O	lis tRNZ , comple	A-Ala, ete cd	phosphas, and l	atidyl RecA (glycer recA)	cophosphate gene,partial cds.]
ORF Name	NTID	AAID	NT LengthI	<u>AA</u> Length	score	probability
A17503000983_9814213_£3_499	632	4404	471	156	232	1.9e-19
Description pir: [LN:C69419] [AC:C69419] [PN:pho phosphate-binding protein (phoX) hom [DB:pir2] >gp:[GI:g2649219] [LN:AE00 transporter, periplasmic] [GN:AF1356 [DE:Archaeoglobus fulgidus section 9 PID:1052826 percent identity: 25.09;	olog] [0 1010] [2] [OR:A: 7 of 172	CL:sph AC:AE0 rchaeo 2 of t	X prote: 01010:Al globus : he comp.	in] [O E00078 fulgid lete g	R:Arch 2] [PN us] [I enome.	naeoglobus fulgidus] I:phosphate ABC DB:genpept-bct2]] [NT:similar to
ORF Name	NTID	AAID	<u>NT</u> LengthI	AA Length	score	probability
A17503000983_9862675_t2_310	633	4405		44		
Description NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> LengthI	<u>AA</u> ength	score	probability
A17503000983_995967_c2_888 Description	634	4406	747	248	341	5.4e-31
pir:[LN:B70039] [AC:B70039] [PN:hypesubtilis] [DB:pir2] >gp:[GI:e1186096 [GN:yvfS] [FN:unknown] [OR:Bacillus complete genome (section 18 of 21): [DI:complement] >gp:[GI:e313010:g194 protein] [GN:yvfS] [OR:Bacillus subtfragment (88 kb).] [NT:probable permeters]	:g263592 subtilis from 339 5719] [I ilis] [I	21] [Li s] [DB 99551to LN:BSZ: OB:gen	N:BSUB00 :genpept o 360906 94043] pept-bct	018] [2 -bct1 50.] [3 [AC:Z94	AC:Z99] [DE: LE:971 4043] E:B.su	121:AL009126] Bacillus subtilis 27] [RE:97864] [PN:hypothetical btilis genomic DNA
ORF Name	NTID	AAID	<u>NT</u> Length1	AA ength	score	probability
A17503000984_10017151_f3_388	635	4407	132	13		
Description NO-HIT						

ORF Name	NTID	AAID NT AA score probability
A17503000984_10039050_f1_17	636	4408 165 54
Description		
NO-HIT		
ORF Name	NTID	AAID <u>NT</u> AA score probability
A17503000984_10193760_f1_89	637	4409 165 54
Description		
NO-HIT		
		NTT 2.2
ORF Name	NTID	AAID <u>NT AA</u> LengthLength score probability
A17503000984_10553125_f3_371	638	4410 810 269 335 1.6e-31
Description		
<pre>gp:[GI:g211700] [LN:CHKCX] [AC:M1349 [SR:Chicken red blood cell DNA, clon [DE:Chicken type X collagen gene.] [</pre>	e pYN92	E1; and embryo chondrocyte] [DB:genpept-vrt]
ORF Name	NTID	AAID Langth Langth Score probability
AT7503000984_10665903_c1_491	639	Length Length
51.6 KD PROTEIN IN TREA-PTH INTERGEN >pir: [LN:C64866] [AC:C64866] [PN:tr [CL:phosphotransferase system phosph [OR:Escherichia coli] [DB:pir2] >gp: [AC:D90754:AB001340] [PN:Hypothetica [OR:Escherichia coli] [SR:Escherichia [DB:genpept-bct1] [DE:Escherichia co [NT:ORF_ID:o245#7; similar to SwissP >gp:[GI:g1787448] [LN:AE000218] [AC: [GN:ycgC] [FN:putative transport; No [DB:genpept-bct2] [DE:Escherichia co completegenome.] [NT:f473; 100 pct i [RE:4520] [DI:complement]	IC REGIO ehalase ohistid [GI:d10] l prote a coli(s li genor rot Acco AE00021s t class li K-12 dentica	ine-containing protein homology] 37041:g4062781] [LN:D90754] in in treA 5'region .] [GN:ycgC] strain:K12) DNA, clone:Kohara clone #245] mic DNA. (26.8 - 27.1 min).] ession] [LE:9793] [RE:11214] [DI:complement] 8:U00096] [PN:putative PTS system enzyme I] ified] [OR:Escherichia coli] MG1655 section 108 of 400 of the 1 to fragment YCGC_ECOLI] [LE:3099]
ORF Name A17503000984_10828312_f1_104	NTID 640	AAID <u>NT AA</u> LengthLength score probability [4412 1617 538 272 1.2e-20
Description	لـــــــا	
pir:[LN:D69796] [AC:D69796] [PN:two [GN:yesM] [OR:Bacillus subtilis] [D	B:pir2] nknown] e (sect:	

ORF Name	NTID AAID NT AA score probability
A17503000984_1182765_c3_673	641 4413 7215 2404 2909 0.0
Description	
[OR:Bacillus licheniformis] [DB:genp synthetase operon, completesequence;	F007865] [PN:bacitracin synthetase 3] [GN:bacC] ept-bct2] [DE:Bacillus licheniformis bacitracin BacS (bacS), BcrA (bcrA), BcrB (bcrB), and BcrC ide synthetase; BA3; BacC] [LE:25258] [RE:44337]
ORF Name	NTID AAID NT AA score probability
AI7503000984_11832518_c2_558	642 4414 144 47
Description NO-HIT	
ORF Name	NTID AAID NT AA score probability
A17503000984_11895058_c1_444	643 4415 939 312 164 5.1e-10
[DB:pir2] >gp:[GI:g43338] [LN:EFSPRE	<pre>ine proteinase homolog] [OR:Enterococcus faecalis] G] [AC:Z12296] [PN:Staphylococcal serine proteinase s faecalis] [DB:genpept-bct1] [DE:E.faecalis sprE .] [LE:91] [RE:945] [DI:direct]</pre>
ORF Name	NTID AAID NT AA score probability
A17503000984_12554627_c3_625 Description	644 4416 216 71
NO-HIT	
ORF Name	NTID AAID NT AA score probability
AI7503000984_1367200_f2_264	645 4417 624 207 104 0.00095
Description sp:[LN:VS10_ROTBS] [AC:P34718] [GN:S [DE:MINOR OUTER CAPSID PROTEIN (NS26	10] [OR:BOVINE ROTAVIRUS] [SR:GROUP C / SHINTOKU,])] [SP:P34718] [DB:swissprot]
ORF Name	NTID AAID NT AA score probability
A17503000984_1367202_c1_489	646 4418 963 320 516 1.1e-60
Description	
[DB:pir2] >gp:[GI:d1037043:g4062783] protein] [OR:Escherichia coli] [SR:E #245] [DB:genpept-bct1] [DE:Escheric [NT:ORF_ID:o246#2; similar to PIR Ac [DI:complement] >gp:[GI:g1787450] [Li	othetical protein b1200] [OR:Escherichia coli] [LN:D90754] [AC:D90754:AB001340] [PN:Hypothetical scherichia coli(strain:K12) DNA, clone:Kohara clone hia coli genomic DNA. (26.8 - 27.1 min).] cession Number] [LE:11865] [RE:12965] N:AE000218] [AC:AE000218:U00096] [PN:putative] [GN:b1200] [FN:putative enzyme; Not classified]

400 of the completegenome.] [NT:f366; 35 pct identical (32 gaps) to 355 residues]

[LE:5171] [RE:6271] [DI:complement]

ORF Name	NTID	AAID NT AA score probability
A17503000984_1367340_c1_424	647	4419 843 280 250 9.9e-31
Description		
	ept-bct	:X89237] [PN:oligopeptidepermease] [GN:oppD] 1] [DE:S.pyogenes DNA for oppA, oppB, oppC, 6924] [DI:direct]
ORF Name	NTID	AAID NT AA score probability
A17503000984_13707008_f3_321	648	4420 231 76
Description		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
AI7503000984_1385962_c3_666	649	4421 984 327 1016 1.6e-102
SYNTHASE, (BIOTIN SYNTHETASE)] [SP:FINE [PN:biotin synthetase bioB] [GN:bioE] [DB:pir2] >gp:[GI:g1277029] [LN:BSUSE] [FN:biotin pathway] [OR:Bacillus subbiotin biosynthetic operon genes, completed [DI:direct] >gp:[GI:e1185893:g263550] synthetase] [GN:bioB] [FN:biotin bioE] [EC:2.8.1] [DE:Bacillus subtilis of 3213410.] [SP:P53557] [LE:91793] [RESTRICT [RESTRICT [SP:P53557]] [RESTRICT [SP:P53557]]	253557] 3] [CL:] 3 [CL:] 3 [CL:] 4 [CL:] 5 [CL:] 5 [CL:] 6 [CL:] 6 [CL:] 6 [CL:] 6 [CL:] 7 [C	BSUB0016] [AC:Z99119:AL009126] [PN:biotin is] [OR:Bacillus subtilis] [DB:genpept-bct1] genome (section 16 of 21): from 2997771to [DI:complement] >gp:[GI:g2293187] hase] [GN:bioB] [OR:Bacillus subtilis]
ORF Name	NTID	AAID NT AA score probability
AI7503000984_14492327_c1_486	650	4422 429 142 495 2.6e-47
RESISTANCE PROTEIN] [SP:Q03377] [DB: [PN:fosfomycin resistance protein B] [OR:Staphylococcus epidermidis] [DB:	swisspro [CL:fos pir2] >q nidis] []	sfomycin resistance protein] gp:[GI:g46982] [LN:SEFOSB] [AC:X54227] DB:genpept-bct1] [DE:S.epidermidis plasmid
ORF Name	NTID	AAID NT AA score probability
A17503000984_14510962_c1_446	651	AAID LengthLength score probability [4423 138 45
Description		

ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000984_14895212_c3_639	652	4424	822 273 980 1.1e-98
Description			
	ylococc de trans utative brane po pp-1D),	us aur sporte membr ermeas and o	reus] [DB:genpept-bct2] er putative substratebinding domain ranepermease domain (opp-1B),
ORF Name	NTID	AAID	NT AA score probability
AI7503000984_157625_c1_428	653	4425	930 309 111 0.00054
Description			
<pre>gp:[GI:e1407888:g4493994] [LN:PFMAL3] falciparum] [SR:malaria parasite P. falciparum MAL3P7, complete sequence (PFC1065w),] [LE:188885] [RE:191470]</pre>	falcipa: .] [NT:]	rum] [] predic	[DB:genpept-inv1] [DE:Plasmodium
ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000984_162550_c3_650	654	4426	198 65 49 0.035
Description			
[AC:F69938] [PN:hypothetical protein >gp:[GI:g1146245] [LN:BACYPIA] [AC:L4 subtilis] [DB:genpept-bct1] [DE:Bacil	n ypmB] 47709] llus sul panBCD ene, cor 683:g263 subtilis	[GN:y] [GN:ypo btilis genes mpleted 34656] s] [DB	(clone YAC15-6B) ypiABF genes, qcrABC, dinG gene, ypmB gene,aspB gene, asnS cds's.] [NT:putative] [LE:18616] [LN:BSUB0012] [AC:Z99115:AL009126] cgenpept-bct1] [DE:Bacillus subtilis
ORF Name	NTID	AAID	NT AA probability
AI7503000984_162578_c1_425	655	4427	984 327 338 1.1e-30
Description pir: [LN:A69867] [AC:A69867] [PN:cons [OR:Bacillus subtilis] [DB:pir2] >gp: [AC:AJ222587] [PN:YkuT protein] [GN:y [DE:Bacillus subtilis 29kB DNA fragme [RE:26383] [DI:complement] >gp:[GI:el [AC:Z99111:AL009126] [GN:ykuT] [FN:ur [DE:Bacillus subtilis complete genome [NT:similar to hypothetical proteins]	:[GI:e11 ykuT] [Cent from 1185011: nknown] e (secti	181925 DR:Bac: m ykwC :g2633 [OR:Ba ion 8 c	:g2632241] [LN:BS16829KB] illus subtilis] [DB:genpept-bct1] gene to cse15 gene.] [LE:25580] 792] [LN:BSUB0008] acillus subtilis] [DB:genpept-bct1] of 21): from 1394791to 1603020.]
ORF Name AI7503000984_16486075_f2_138	NTID	AAID [4428	NT AA LengthLength score probability
Description	لــــــــــــــــــــــــــــــــــــــ		التعميا
NO-HIT			

ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	score	probability
AI7503000984_165902_c3_674	657	4429		208	185	1.8e-14
Description sp:[LN:LP14_BACSU] [AC:P39144] [GN:Li ANTIBIOTICS ITURIN A AND SURFACTIN B: >pir:[LN:I39875] [AC:I39875] [PN:sic sfp:lipopeptide antibiotics iturin A [GN:lpa-14:sfb:sfp:sfp(0)] [CL:side: [OR:Bacillus subtilis] [DB:pir2] >gp [PN:lipopeptide antibiotics iturin A subtilis (strain:RB14) DNA] [DB:genpellipopeptide antibiotics iturin A.]	IOSYNTHI deropho: :surfact rophore :[GI:d10] [GN:l] ept-bct	ESIS Pire bios tin pro biosyn 005421 pa-14] [DE	ROTEIN] synthes oductio nthesis :g47391 [OR:Ba :B. sub	[SP:Fis reg n prot regul 6] [LN cillus	239144] rulator ein] atory :BACLE subti	[DB:swissprot] Ty protein protein sfp] PA14] [AC:D21876] Lis] [SR:Bacillus
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000984_187561_c1_470	658	4430	879	292	0	
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
AI7503000984_194010_c3_607	659	4431	1512	503	1381	3.4e-141
<pre>gp:[GI:d1039113:g4514332] [LN:AB01336 [SR:Bacillus halodurans (strain:C-125 C-125 yesT and comEC genes, partial a [DI:direct]</pre>	5) DNA]	[DB:ge	enpept-	bct1]	[DE:Ba	cillus halodurans
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000984_19688401_c2_516	660	4432	126	41		
Description NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Lengthl	<u>AA</u> Length	score	probability
AI7503000984_19773387_£3_363	661	4433	1248	415	746	6.6e-74
Description sp:[LN:STPA_STAAU] [AC:P81297] [OR:ST [DE:STAPHOPAIN,] [SP:P81297] [DB:swis		coccus	AUREUS] [EC:	3.4.22]
ORF Name	NTID	AAID	<u>NT</u> Length]	<u>AA</u> Length	score	probability
AI7503000984_20585963_f3_396	662	4434	897	298	657	1.8e-64
Description [17]				_		
gp:[GI:g929972] [LN:BAU30714] [AC:U30 [DB:genpept-bct1] [DE:Bacillus anthra invertedrepeat element (WeyAR) border complete cds.] [NT:ORFB; similar to F [RE:1336] [DI:direct] >gp:[GI:g929975 anthracis] [SR:plasmid pXO1] [DB:genpelasmid pXO1] left inverted repeateler ORFB andtruncated ORFA genes, complete element ORFB;] [LE:458] [RE:1282] [DI [AC:AF065404] [PN:pXO1-96] [OR:Bacillanthracis virulence plasmid PXO1, complete com	acis Wey cing the 3. anthr 5] [LN:F pept-bct ment (St ce cds.] I:direct Lus anth	ybridge toxir cacis S BAU3071 [DE cerneL) [NT:0] ygp:	e A tox. n-encod. SterneL. 5] [AC ::Bacil: borde: DRFB; s: [GI:g4	in pla ing re eleme :U3071 lus an ring t imilar 894312 enpept	smid p gion, nt ORF 5] [OR thraci he tox to B.] [LN: -bct2]	XO1 right ORFAand ORFB genes, B;] [LE:512] :Bacillus s Sterne toxin in-encoding region, anthracis WeyAR AF065404] [DE:Bacillus

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000984_20704012_t1_106	663	4435	7240	79	: }	
Description			لــــا		J	
NO-HIT						
					200 00.000 00.000	and the second s
ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	score	probability
A17503000984_20979688_c2_511	664	4436	786	261	369	5.9e-34
Description sp:[LN:NIKC_ECOLI] [AC:P33592] [GN:N SYSTEM PERMEASE PROTEIN NIKC] [SP:P3 [AC:S39596:S47697:A65145] [PN:nikC protein oppB] [OR:Escherichia coli] [PN:NikC] [GN:nikC] [OR:Escherichia nik locus.] [SP:P33592] [LE:2942] [R [AC:U00039] [GN:nikC] [OR:Escherichi strain K-12) (library: lambda] [DB:g 76.0 to 81.5 minutes.] [LE:30444] [R [LN:AE000423] [AC:AE000423:U00096] [[FN:transport; Transport of small mo [DE:Escherichia coli K-12 MG1655 sec [LE:7496] [RE:8329] [DI:direct]	3592] [] protein [DB:pir: coli] [] E:3775] a coli] enpept-] E:31277] PN:trans lecules	DB:swi] [GN: 2] >gp DB:gen [DI:d [SR:E bctl]] [DI: sport :] [OR	ssprot] nikC] :[GI:g5 pept-bo irect] scheric [DE:E. direct] of nick :Escher	>pir: [CL:ol 581141] ctl] [D >gp:[G chia co coli c >gp:[cel, me	[LN:S3 igopep [LN:E E:E.co I:g912 li (su hromos GI:g17 mbrane coli]	tide permease CNIK] [AC:X73143] li DNA sequence of 461] [LN:ECOUW76] b_strain MG1655, omal region from 89889] protein] [GN:nikC] [DB:genpept-bct2]
ORF Name	NTID	AAID	NT	<u>AA</u>	score	probability
A17503000984_212827_f3_295 Description	665	4437	Length 1482	Length 493		2.6e-40
pir:[LN:G70006] [AC:G70006] [PN:mul [OR:Bacillus subtilis] [DB:pir2] >gp [AC:Z99119:AL009126] [GN:yubD] [FN:ul [DE:Bacillus subtilis complete genom [NT:similar to multidrug resistance]	:[GI:e1: nknown] e (sect:	185986 [OR:B ion 16	g26355: acillus of 21)	97] [L s subti : from	N:BSUB lis] [29977	0016] DB:genpept-bct1] 71to 3213410.]
ORF Name A17503000984_21588287_t2_271	NTID 666	<u>AAID</u>	<u>NT</u> Length	AA Length 41	score	probability
Description NO-HIT			ليساد			
ORF Name AI7503000984_21907016_f1_123	<u>NTID</u> 667	<u>AAID</u> 4439	NT Length	AA Length 43	score	probability
Description						

ORF Name	NTID AAID LengthLength score probability
AI7503000984_22042337_c2_588	668 4440 1434 477 981 8.3e-99
Description	
[CL:lincomycin-resistance protein lm >gp:[GI:e1182351:g2632685] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-b of 21): from 402751 to611850.] [NT [LE:32866] [RE:34284] [DI:complement [AC:D50453] [PN:homologue of multidr subtilis] [SR:Bacillus subtilis (str	O03] [AC:Z99106:AL009126] [GN:ycnB] [FN:unknown] ctl] [DE:Bacillus subtilis complete genome (section :similar to multidrug resistance protein] >gp:[GI:d1009651:g1805454] [LN:D50453] ug resistance protein B,] [GN:ycnB] [OR:Bacillus ain:168 trpC2) DNA] [DB:genpept-bctl] [DE:Bacillus containing theamyE-srfA region, complete cds.]
ORF Name	NTID AAID NT AA score probability
A17503000984_22323413_c2_518	669 4441 144 47
Description NO-HIT	
ORF Name	NTID AAID NT AA score probability
AI7503000984_22464127_f2_191	670 4442 153 50
Description	
NO-HIT	
ORF Name	NTID AAID <u>NT AA</u> score probability
A17503000984_22537818_c2_513	671 4443 150 49
Description NO-HIT	
100-1111	
ORF Name	NTID AAID LengthLength score probability
AI7503000984_22664140_t3_355	672 4444 1395 464 191 1.0e-11
iron transport protein B:translation fulgidus] [DB:pir2] >gp:[GI:g2650395 (II) transporter (feoB-1)] [GN:AF024 [DE:Archaeoglobus fulgidus section 18	n (II) transporter (feoB-1) homolog] [CL:ferrous elongation factor Tu homology] [OR:Archaeoglobus] [LN:AE001089] [AC:AE001089:AE000782] [PN:iron 6] [OR:Archaeoglobus fulgidus] [DB:genpept-bct2] 8 of 172 of the complete genome.] [NT:similar to ent] [LE:10039] [RE:11958] [DI:complement]
ORF Name	NTID AAID <u>NT AA</u> score probability
A17503000984_22664550_c2_512	673 4445 663 220 275 5.4e-24
ATP-binding cassette proteins: ATP-binding protein of nickel tract case case case case case case case case	[7699] [PN:nikE protein] [GN:nikE] [CL:unassigned inding cassette homology] [OR:Escherichia coli] [76] [AC:U00039] [GN:nikE] [OR:Escherichia coli] [555, strain K-12) (library: lambda] [Somal region from 76.0 to 81.5 minutes.] [LE:32038] [Somal region from 76.0 to 81.5 minutes.] [S

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
AI7503000984_22853432_c1_448	674	4446	384	127	152	2.4e-10
Description Sp:[LN:INVO_PIG] [AC:P18175] [GN:IVL [SP:P18175] [DB:swissprot] >pir:[LN: [CL:involucrin] [OR:Sus scrofa domes >gp:[GI:g164523] [LN:PIGINVOLA] [AC: skin keratinocyte DNA] [DB:genpept-m [NT:involucrin] [LE:1] [RE:1044] [DI	I46592] tica] [M34441] am] [DE	AC:I4 SR:, do OR:Su Pig in:	16592] omestic us scro	[PN:i pig] ofa] [S	nvolud [DB:pi R:Pig	erin] ir2] (Yorkshire) adult
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
17503000984_23444425_c1_426	675	4447	846	281	362	3.2e-33
pir:[LN:B69834] [AC:B69834] [PN:con [OR:Bacillus subtilis] [DB:pir2] >gp [AC:Z99109:AL009126] [GN:yhjK] [FN:u [DE:Bacillus subtilis complete genom [NT:similar to hypothetical proteins >gp:[GI:e324984:g2226183] [LN:BSY140 [OR:Bacillus subtilis] [DB:genpept-b 92 degrees: regionbetween comK and a hypothetical] [LE:9722] [RE:10582] [:[GI:e1 nknown] e (sect] [LE:1 81] [AC ct1] [D ddAB.]	183056; [OR:Baion 6 c 27427] :Y14081 E:Bacil [NT:Sin	g26333 acillus of 21): [RE:12 l] [PN: llus su	390] [Lis subti from 28287] hypoth	N:BSUE lis] 999501 [DI:co etica] chron	B0006] [DB:genpept-bct1] L to1209940.] complement] L protein] [GN:yhjK] cosomal DNA, region
DRF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
AI7503000984_23470290_c2_554	676	4448		304	1205	1.5e-122
Description Gescription Gescr	ccus au tive su e domai p-1C), ansport	reus] bstrate n (opp- oligope erputat	[DB:gen ebindin -1B), c eptide :ive AT	npept-b ng doma bligope transp Pase d	ct2] in (op ptide orterp omain	DE:Staphylococcus pp-1A), oligopeptide transporter putative ATPase
ORF Name A17503000984 235837 cl 471	NTID	<u>AAID</u>	<u>NT</u> Length	<u>AA</u> Length	score	probability
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
17503000984_23601510_c1_488	678	4450	1137	378	911	2.2e-91
Description Sp: [LN:GLDA_BACST] [AC:P32816] [GN:GET:[LN:GLDA_BACST]] [AC:P32816] [GN:GET:[LN:JQ1474]] [AC:JQ1474:S38514] [CL:glycerol dehydrogenase:lactaldehystearothermophilus] [EC:1.1.1.6] [DB:PN:glycerol dehydrogenase] [GN:gld] Stearothermophilus (sub_species nondefice:1.1.1.6] [DE:Bacillus stearothermophilus]	ASE, (GI [PN:gly yde redu :pir2] : [OR:Badiastation	LDH)] [ycerol uctase >gp:[GI cillus cus) (1	SP:P32 dehydr homolo :g1429 stearo ibrary	816] [] ogenase gy] [O] 78] [LI othermon	OB:swi e,] [G R:Baci N:BACG philus genpep	ssprot] N:gldA] llus LDA] [AC:M65289]] [SR:Bacillus ot-bct1]

ORF Name	NTID A	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000984_23652218_c1_459	679	4451	168	55	83	0.0012
Description sp:[LN:GGI3_STAHA] [AC:P11699] [OR:S PROTEIN 3 (GONOCOCCAL GROWTH INHIBIT [AC:S00601] [PN:antibacterial prote [CL:Staphylococcus haemolyticus anti [DB:pir1]	OR 3)] [S in 3:gond	SP:P1	1699] al grov	[DB:swis vth inhi	ssprot ibitor] >pir:[LN:BXSA3]
ORF Name [A17503000984 23860307 c3 641]		AAID	<u>NT</u> Length	Length-	score	probability
Description			لتتتا			
gp:[GI:g3800823] [LN:AF076683] [AC:A [DB:genpept-bct2] [DE:Staphylococcus substratebinding domain (opp-1A), ol domain (opp-1B), oligopeptide transpoligopeptide transporterputative ATP transporterputative ATPase domain (o [NT:orfX] [LE:4904] [RE:6097] [DI:di	aureus o igopeptio orter put ase domai pp-1F) ge	oligon de tra cativa in (on	peptide ansport emembra pp-1D),	e transp er puta ne perm and ol	porter ative nease Ligope	putative membranepermease domain (opp-1C), ptide
ORF Name	NTID E	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
AI7503000984_24000275_f3_303	681 4	1453	1026			2.5e-104
Description sp:[LN:OTCC_HAEIN] [AC:P44770] [GN:A [EC:2.1.3.3] [DE:ORNITHINE CARBAMOYL [DB:swissprot] >pir:[LN:H64079] [AC: [CL:ornithine carbamoyltransferase: [OR:Haemophilus influenzae] [EC:2.1. [AC:U32741:L42023] [PN:ornithine car [OR:Haemophilus influenzae Rd] [DB:gr 56 of 163 of the complete genome.] [Second Carbana (LE:3467] [RE:4471] [DI:complete carbana (LE:3467] [RE:4471] [RE:4471] [DI:complete carbana (LE:3467] [RE:4471] [R	TRANSFERA H64079] aspartate 3.3] [DB: bamoyltra enpept-bo NT:simila	ASE, ([PN:c e/orni pir2] ansfer	CATABOL ornithi ithine >gp:[case (a [DE:Hae	IC, (OI ne carb carbamo [GI:g157 [rcB)] [CCASE) camoyl cyltra 73585] [GN:HI is inf] [SP:P44770] transferase,] nsferase homology] [LN:U32741] 0596] luenzae Rd section
ORF Name	NTID A	AAID	<u>NT</u> Length	<u>AA</u> Length ^s	score	probability
A17503000984_24095387_c1_438 Description NO-HIT	682 4	1454	129	42		
ORF Name A17503000984_24105393_c2_562 Description		AAID 1455	NT Length	Length -	711 [probability 3.4e-70
gp:[GI:e1456529:g4914622] [LN:LMAJ96] activating enzyme] [GN:pflC] [OR:List monocytogenes pflC, orfA, lltB and omutans PflC] [LE:149] [RE:895] [DI:di	teria mon rfC genes	ocyto	genes]	[DB:ge	npept	-bct1] [DE:Listeria

		NT AA
ORF Name	NTID	AAID LengthLength score probability
A17503000984_24225375_c2_533	684	4456 213 70 97 3.9e-05
Description		
<pre>beta 2 genes, complete cds.] [NT:PSM >gp:[GI:g3212080] [LN:AF068633] [AC: [FN:inflammatory protein] [OR:Staphy</pre>	rlococcu ol solub ol beta 1 AF06863 rlococcu ol solub	s epidermidis] [DB:genpept-bct2] le modulin beta 1 and phenolsoluble modulin] [LE:669] [RE:803] [DI:direct] 3] [PN:phenol soluble modulin beta 2] s epidermidis] [DB:genpept-bct2] le modulin beta 1 and phenolsoluble modulin
ORF Name	NTID	AAID NT AA score probability
A17503000984_24257881_f1_68	685	4457 138 45
Description		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000984_24266041_t1_22	686	4458 129 42
Description		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000984_24391678_f1_120	687	Length Length Floor Flor
Description		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000984_24407677_t3_293	688	4460 234 77
Description		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000984_24650300_c1_476	689	4461 1365 454 1037 9.6e-105
Description		
<pre>>pir:[LN:S60180] [AC:S60180] [PN:br [CL:branched-chain amino acid transp [OR:Lactobacillus delbrueckii] [DB:p</pre>	ID UPTAN anched-c ort syst ir2] >gr	KE CARRIER)] [SP:P54104] [DB:swissprot] chain amino acid carrier brnQ] [GN:brnQ] tem II carrier protein braZ] p:[GI:g732813] [LN:LDBRNQGN] [AC:Z48676]
		brnQ] [FN:transport of branched-chain amino [DB:genpept-bct1] [DE:L.delbrueckii brnQ
		[SP:P54104] [LE:611] [RE:1951] [DI:direct]

ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	score	probability
A17503000984_24664012_t2_199	690	4462	453	150	82	0.0052
Description gp:[GI:d1025730:g2879910] [LN:D85752 [SR:Enterococcus faecalis plasmid:pP plasmid pPD1 bacA, bacB, bacC, bacD, cds.] [LE:3977] [RE:4324] [DI:direct	D1 DNA] bacE,ba	[DB:g	enpept-	-bct1]	[DE:E	nterococcus faecalis
ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	score	probability
A17503000984_24711588_c2_589	691	4463	225	74	l	
Description NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000984_24884688_c2_509 Description	692	4464	528	175	216	5.6e-17
gp:[GI:g3800818] [LN:AF076683] [AC:A substrate] [GN:opp-1A] [OR:Staphyloc aureus oligopeptide transporter puta transporter putative membranepermeas putativemembrane permease domain (oppdomain (opp-1D), and oligopeptide transporter cds; and unknowngene.] [LE:	occus antive sul e domain p-1C), d ansporte	ureus] ostrato n (oppoligopo erputa	(DB:ge ebindir -1B), c eptide tive AT	enpept- ng doma oligope transp	bct2] in (op ptide orterp	[DE:Staphylococcus pp-1A), oligopeptide transporter putative ATPase
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000984_24886552_c3_635	693	4465	792	263	603	9.4e-59
Description gp:[GI:d1037145:g4062842] [LN:AB0090 dehydrogenase] [OR:Brevibacterium sa DNA] [DB:genpept-bct1] [DE:Brevibacte L-2.3-butanedioldehydrogenase, comple	ccharoly erium sa	yticum; acchar	[SR:E	Breviba um gene	cterion	um saccharolyticum
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000984_25429665_c1_496	694	4466	1338	445	267	4.3e-21
Description pir: [LN:S58131] [AC:S58131] [PN:intellectis] [DB:pir2] >gp: [GI:g1052754] protein] [GN:lmrP] [OR:Lactococcus lagene.] [LE:634] [RE:1860] [DI:direct	[LN:LLLNactis]	MRP] [A	AC:X897	79] [P	N:Lmr	integral membrane
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	<u>score</u>	probability
A17503000984_25476378_f1_13	695	4467	159	52		
Description NO-HIT						
ORF Name A17503000984_25894687_c3_624	NTID 696	<u>AAID</u> 4468	NT Length	AA Length 80	score	probability
Description NO-HIT						

ORF Name	NTID	AAID NT AA score probability	
AI7503000984_26182681_c3_595	697	[4469 [528] [175]	
Description			
NO-HIT			
	terne mende en mende frestante desse	NT AA	
ORF Name	NTID	AAID LengthLength score probability	
AI7503000984_26208450_f1_24	698	[4470 165 54	
Description			
NO-HIT			
ORF Name	NTID	AAID NT AA score probability LengthLength	
A17503000984_26229678_f1_27	699	4471 177 158	
Description			
NO-HIT			
		NT AA	
ORF Name	NTID	AAID LengthLength score probability	
AI7503000984_26265641_c2_543	700	4472 [147] 48	
Description			
NO-HIT			
ORF Name	NTID	AAID NT AA score probability	
MT7507000004 26267125 at 500			
A17503000984_26367135_c1_508	701	4473 2079 692 1496 2.2e-153	
Description	701	4473 2079 692 1496 2.2e-153	
Description sp:[LN:LIP_STAEP] [AC:Q02510] [GN:GE	HC] [OR	STAPHYLOCOCCUS EPIDERMIDIS] [EC:3.1.1.3]	
Description sp:[LN:LIP_STAEP] [AC:Q02510] [GN:GE [DE:LIPASE PRECURSOR, (GLYCEROL ESTE	HC] [OR R HYDRO	STAPHYLOCOCCUS EPIDERMIDIS] [EC:3.1.1.3] ASE)] [SP:Q02510] [DB:swissprot]	
Description sp:[LN:LIP_STAEP] [AC:Q02510] [GN:GE [DE:LIPASE PRECURSOR, (GLYCEROL ESTE >pir:[LN:A47705] [AC:A47705] [PN:tr	HC] [OR R HYDROI	STAPHYLOCOCCUS EPIDERMIDIS] [EC:3.1.1.3] ASE)] [SP:Q02510] [DB:swissprot] cerol lipase,] [CL:Staphylococcus	
Description sp:[LN:LIP_STAEP] [AC:Q02510] [GN:GE [DE:LIPASE PRECURSOR, (GLYCEROL ESTE >pir:[LN:A47705] [AC:A47705] [PN:tr triacylglycerol lipase] [OR:Staphylo >gp:[GI:g153022] [LN:STAGEHC] [AC:M9	HC] [OR R HYDROI iacylgly coccus e	STAPHYLOCOCCUS EPIDERMIDIS] [EC:3.1.1.3] ASE)] [SP:Q02510] [DB:swissprot] cerol lipase,] [CL:Staphylococcus pidermidis] [EC:3.1.1.3] [DB:pir2] N:lipase] [GN:gehC] [OR:Staphylococcus	
Description sp:[LN:LIP_STAEP] [AC:Q02510] [GN:GE [DE:LIPASE PRECURSOR, (GLYCEROL ESTE >pir:[LN:A47705] [AC:A47705] [PN:tr triacylglycerol lipase] [OR:Staphylo >gp:[GI:g153022] [LN:STAGEHC] [AC:M9 epidermidis] [SR:Staphylococcus epid	HC] [OR R HYDROI (COCCUS 6 5577] [1] ermidis	STAPHYLOCOCCUS EPIDERMIDIS] [EC:3.1.1.3] ASE)] [SP:Q02510] [DB:swissprot] cerol lipase,] [CL:Staphylococcus pidermidis] [EC:3.1.1.3] [DB:pir2] N:lipase] [GN:gehC] [OR:Staphylococcus (strain 9) DNA] [DB:genpept-bct1]	
Description sp:[LN:LIP_STAEP] [AC:Q02510] [GN:GE [DE:LIPASE PRECURSOR, (GLYCEROL ESTE >pir:[LN:A47705] [AC:A47705] [PN:tr triacylglycerol lipase] [OR:Staphylo >gp:[GI:g153022] [LN:STAGEHC] [AC:M9 epidermidis] [SR:Staphylococcus epid [DE:Staphylococcus epidermidis lipas	HC] [OR R HYDROI (COCCUS 6 5577] [1] ermidis	STAPHYLOCOCCUS EPIDERMIDIS] [EC:3.1.1.3] ASE)] [SP:Q02510] [DB:swissprot] cerol lipase,] [CL:Staphylococcus pidermidis] [EC:3.1.1.3] [DB:pir2] N:lipase] [GN:gehC] [OR:Staphylococcus	1]
Description sp:[LN:LIP_STAEP] [AC:Q02510] [GN:GE [DE:LIPASE PRECURSOR, (GLYCEROL ESTE >pir:[LN:A47705] [AC:A47705] [PN:tr triacylglycerol lipase] [OR:Staphylo >gp:[GI:g153022] [LN:STAGEHC] [AC:M9 epidermidis] [SR:Staphylococcus epid	HC] [OR R HYDROI (COCCUS 6 5577] [1] ermidis	STAPHYLOCOCCUS EPIDERMIDIS] [EC:3.1.1.3] ASE)] [SP:Q02510] [DB:swissprot] cerol lipase,] [CL:Staphylococcus pidermidis] [EC:3.1.1.3] [DB:pir2] N:lipase] [GN:gehC] [OR:Staphylococcus (strain 9) DNA] [DB:genpept-bct1] gene, complete cds.] [NT:GTG start codon	1]
Description sp:[LN:LIP_STAEP] [AC:Q02510] [GN:GE [DE:LIPASE PRECURSOR, (GLYCEROL ESTE >pir:[LN:A47705] [AC:A47705] [PN:tr triacylglycerol lipase] [OR:Staphylo >gp:[GI:g153022] [LN:STAGEHC] [AC:M9 epidermidis] [SR:Staphylococcus epid [DE:Staphylococcus epidermidis lipas	HC] [OR R HYDROI (COCCUS 6 5577] [1] ermidis	STAPHYLOCOCCUS EPIDERMIDIS] [EC:3.1.1.3] ASE)] [SP:Q02510] [DB:swissprot] cerol lipase,] [CL:Staphylococcus pidermidis] [EC:3.1.1.3] [DB:pir2] N:lipase] [GN:gehC] [OR:Staphylococcus (strain 9) DNA] [DB:genpept-bct1]	1]
Description sp:[LN:LIP_STAEP] [AC:Q02510] [GN:GE [DE:LIPASE PRECURSOR, (GLYCEROL ESTE >pir:[LN:A47705] [AC:A47705] [PN:tr triacylglycerol lipase] [OR:Staphylo >gp:[GI:g153022] [LN:STAGEHC] [AC:M9 epidermidis] [SR:Staphylococcus epid [DE:Staphylococcus epidermidis lipas [LE:121] [RE:2187] [DI:direct]	HC] [OR R HYDROD liacylgly coccus 6 5577] [1 ermidis e (gehC)	STAPHYLOCOCCUS EPIDERMIDIS] [EC:3.1.1.3] ASE)] [SP:Q02510] [DB:swissprot] cerol lipase,] [CL:Staphylococcus pidermidis] [EC:3.1.1.3] [DB:pir2] N:lipase] [GN:gehC] [OR:Staphylococcus (strain 9) DNA] [DB:genpept-bct1] gene, complete cds.] [NT:GTG start codon	1]
Description sp:[LN:LIP_STAEP] [AC:Q02510] [GN:GE [DE:LIPASE PRECURSOR, (GLYCEROL ESTE >pir:[LN:A47705] [AC:A47705] [PN:tr triacylglycerol lipase] [OR:Staphylo >gp:[GI:g153022] [LN:STAGEHC] [AC:M9 epidermidis] [SR:Staphylococcus epid [DE:Staphylococcus epidermidis lipas [LE:121] [RE:2187] [DI:direct] ORF Name	HC] [OR R HYDROD RICOCCUS 6 S577] [1] ermidis e (gehC)	STAPHYLOCOCCUS EPIDERMIDIS] [EC:3.1.1.3] [ASE)] [SP:Q02510] [DB:swissprot] [Cerol lipase,] [CL:Staphylococcus [pidermidis] [EC:3.1.1.3] [DB:pir2] [N:lipase] [GN:gehC] [OR:Staphylococcus [(strain 9) DNA] [DB:genpept-bct1] [gene, complete cds.] [NT:GTG start codon [AAID NT AA [LengthLength] score probability	1]
Description sp:[LN:LIP_STAEP] [AC:Q02510] [GN:GE [DE:LIPASE PRECURSOR, (GLYCEROL ESTE >pir:[LN:A47705] [AC:A47705] [PN:tr triacylglycerol lipase] [OR:Staphylo >gp:[GI:g153022] [LN:STAGEHC] [AC:M9 epidermidis] [SR:Staphylococcus epid [DE:Staphylococcus epidermidis lipas [LE:121] [RE:2187] [DI:direct] ORF Name AT7503000984_26369027_f2_260 Description pir:[LN:E69806] [AC:E69806] [PN:con	HC] [OR R HYDROD iacylgly coccus of 5577] [1] ermidis e (gehC) NTID	STAPHYLOCOCCUS EPIDERMIDIS] [EC:3.1.1.3] [ASE)] [SP:Q02510] [DB:swissprot] [Cerol lipase,] [CL:Staphylococcus [pidermidis] [EC:3.1.1.3] [DB:pir2] [N:lipase] [GN:gehC] [OR:Staphylococcus [strain 9) DNA] [DB:genpept-bct1] [gene, complete cds.] [NT:GTG start codon [AAID NT AA LengthLength score probability [4474 984 327 1350 6.5e-138 [Sypothetical protein yfjN] [GN:yfjN]	1]
Description sp:[LN:LIP_STAEP] [AC:Q02510] [GN:GE [DE:LIPASE PRECURSOR, (GLYCEROL ESTE >pir:[LN:A47705] [AC:A47705] [PN:tr triacylglycerol lipase] [OR:Staphylo >gp:[GI:g153022] [LN:STAGEHC] [AC:M9 epidermidis] [SR:Staphylococcus epid [DE:Staphylococcus epidermidis lipas [LE:121] [RE:2187] [DI:direct] ORF Name A17503000984_26369027_f2_260 Description pir:[LN:E69806] [AC:E69806] [PN:con [OR:Bacillus subtilis] [DB:pir2] >gp	HC] [OR R HYDROD iacylgly coccus of 5577] [1] ermidis e (gehC) NTID 702 served h: [GI:ell	STAPHYLOCOCCUS EPIDERMIDIS] [EC:3.1.1.3] [ASE)] [SP:Q02510] [DB:swissprot] [Cerol lipase,] [CL:Staphylococcus [pidermidis] [EC:3.1.1.3] [DB:pir2] [N:lipase] [GN:gehC] [OR:Staphylococcus [strain 9) DNA] [DB:genpept-bct1] [gene, complete cds.] [NT:GTG start codon [AAID NT AA LengthLength Score probability [4474 984 327 1350 6.5e-138 [Sypothetical protein yfjN] [GN:yfjN] [Sypothetical protein yfjN] [GN:yfjN] [Sypothetical protein yfjN] [GN:yfjN]	1]
Description sp:[LN:LIP_STAEP] [AC:Q02510] [GN:GE [DE:LIPASE PRECURSOR, (GLYCEROL ESTE >pir:[LN:A47705] [AC:A47705] [PN:tr triacylglycerol lipase] [OR:Staphylo >gp:[GI:g153022] [LN:STAGEHC] [AC:M9 epidermidis] [SR:Staphylococcus epid [DE:Staphylococcus epidermidis lipas [LE:121] [RE:2187] [DI:direct] ORF Name A17503000984_26369027_f2_260 Description pir:[LN:E69806] [AC:E69806] [PN:con [OR:Bacillus subtilis] [DB:pir2] >gp [AC:Z99108:AL009126] [GN:yfjN] [FN:u	HC] [OR R HYDROD iacylgly coccus of 5577] [I ermidis e (gehC) NTID 702 served large serve	STAPHYLOCOCCUS EPIDERMIDIS] [EC:3.1.1.3] [ASE)] [SP:Q02510] [DB:swissprot] [Cerol lipase,] [CL:Staphylococcus [pidermidis] [EC:3.1.1.3] [DB:pir2] [N:lipase] [GN:gehC] [OR:Staphylococcus (strain 9) DNA] [DB:genpept-bct1] gene, complete cds.] [NT:GTG start codon [AAID NT AA LengthLength Score probability [A474 984 327 1350 6.5e-138 [Sypothetical protein yfjN] [GN:yfjN] [Sypothetical protein yfjN] [GN:yfjN] [Sypothetical subtilis] [DB:genpept-bct1]	.]
Description sp:[LN:LIP_STAEP] [AC:Q02510] [GN:GE [DE:LIPASE PRECURSOR, (GLYCEROL ESTE >pir:[LN:A47705] [AC:A47705] [PN:tr triacylglycerol lipase] [OR:Staphylo >gp:[GI:g153022] [LN:STAGEHC] [AC:M9 epidermidis] [SR:Staphylococcus epid [DE:Staphylococcus epidermidis lipas [LE:121] [RE:2187] [DI:direct] ORF Name AT7503000984_26369027_f2_260 Description pir:[LN:E69806] [AC:E69806] [PN:con [OR:Bacillus subtilis] [DB:pir2] >gp [AC:Z99108:AL009126] [GN:yfjN] [FN:u [DE:Bacillus subtilis complete genom [NT:similar to hypothetical proteins	HC] [OR R HYDROD iacylgly coccus of 5577] [1] ermidis e (gehC) NTID 702 served language in the company of th	STAPHYLOCOCCUS EPIDERMIDIS] [EC:3.1.1.3] [ASE)] [SP:Q02510] [DB:swissprot] [Cerol lipase,] [CL:Staphylococcus [Price pidermidis] [EC:3.1.1.3] [DB:pir2] [N:lipase] [GN:gehC] [OR:Staphylococcus (strain 9) DNA] [DB:genpept-bct1] [gene, complete cds.] [NT:GTG start codon [AAID NT AA LengthLength score probability [A474 984 327 1350 6.5e-138 [Aypothetical protein yfjN] [GN:yfjN] [Sypothetical protein yfjN] [GN:yfjN] [Sypothetical protein yfjN] [DB:genpept-bct1] [OR:Bacillus subtilis] [DB:genpept-bct1] on 5 of 21): from 802821 to1011250.] 112] [RE:74089] [DI:direct]	13
Description sp:[LN:LIP_STAEP] [AC:Q02510] [GN:GE [DE:LIPASE PRECURSOR, (GLYCEROL ESTE >pir:[LN:A47705] [AC:A47705] [PN:tr triacylglycerol lipase] [OR:Staphylo >gp:[GI:g153022] [LN:STAGEHC] [AC:M9 epidermidis] [SR:Staphylococcus epid [DE:Staphylococcus epidermidis lipas [LE:121] [RE:2187] [DI:direct] ORF Name AT7503000984_26369027_f2_260 Description pir:[LN:E69806] [AC:E69806] [PN:con [OR:Bacillus subtilis] [DB:pir2] >gp [AC:Z99108:AL009126] [GN:yfjN] [FN:u [DE:Bacillus subtilis complete genom [NT:similar to hypothetical proteins >gp:[GI:d1025211:g2780398] [LN:D7850]	HC] [OR R HYDROD iacylgly coccus of 5577] [1] ermidis e (gehC) NTID 702 served h :[GI:elinknown] e (section of 100 coccus) e (section of 100 coccus) [LE:739] [AC:I	STAPHYLOCOCCUS EPIDERMIDIS] [EC:3.1.1.3] [ASE)] [SP:Q02510] [DB:swissprot] [Cerol lipase,] [CL:Staphylococcus [pidermidis] [EC:3.1.1.3] [DB:pir2] [N:lipase] [GN:gehC] [OR:Staphylococcus (strain 9) DNA] [DB:genpept-bct1] gene, complete cds.] [NT:GTG start codon [AAID NT AA LengthLength score probability [4474 984 327 1350 6.5e-138 [Cypothetical protein yfjN] [GN:yfjN] [82793:g2633127] [LN:BSUB0005] [OR:Bacillus subtilis] [DB:genpept-bct1] on 5 of 21): from 802821 to1011250.]	1]

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probab	ility
A17503000984_26600936_f2_253	703	4475	513	170	373	2.2e-34	
Description							· · · · · · · · · · · · · · · · · · ·
sp:[LN:Y318_HAEIN] [AC:P43984] [GN:HPROTEIN HI0318] [SP:P43984] [DB:swis [PN:hypothetical protein HI0318] [OR >gp:[GI:g1573288] [LN:U32717] [AC:U3 [GN:HI0318] [OR:Haemophilus influenz Rd section 32 of 163 of the complete GB:AL009126] [LE:4064] [RE:4582] [DI	sprot] :Haemop 2717:L4 ae Rd] genome	>pir:[] hilus : 2023] [DB:ger .] [NT:	EN:B640 influen [PN:con npept-b	06] [A zae] [served ct2] [C:B640 DB:pir hypot DE:Hae	06] 2] hetical mophilus	protein] influenzae
ORF Name	NTID	AAID	<u>NT</u> Lengthl	<u>AA</u> Length	score	probab	ility
AI7503000984_26759430_c3_654	704	4476	978	325	872	2.9e-87	
Description pir: [LN:E69670] [AC:E69670] [PN:gly (osmoprotec) opuCC] [GN:opuCC] [OR: >gp: [GI:e1186069:g2635894] [LN:BSUB0 betaine/carnitine/choline ABC] [GN:o betaine,] [OR:Bacillus subtilis] [DB genome (section 18 of 21): from 3399 [LE:67766] [RE:68677] [DI:complement	Bacillu 018] [A puCC] [:genpep 551to 3	s subti C:Z9912 FN:high t-bct1]	llis] [1 21:AL00 1 affin [DE:B	DB:pir 9126] ity tra acillu	2] [PN:gl anspor s subt	ycine t of gly ilis com	cine plete
ORF Name	NTID	AAID	<u>NT</u> Lengthl	<u>AA</u> Length	score	probab	ility
A17503000984_26760076_f3_305	705	4477	1578	525	1550	4.2e-159	
Description sp:[LN:YFCC_HAEIN] [AC:P44023] [GN:H PROTEIN HI0594] [SP:P44023] [DB:swis [PN:hypothetical protein HI0594] [CL protein HI0594] [OR:Haemophilus infl [AC:U32741:L42023] [PN:conserved hyp [OR:Haemophilus influenzae Rd] [DB:g 56 of 163 of the complete genome.] [[LE:891] [RE:2420] [DI:complement]	sprot] :Haemop uenzae] othetic enpept-	>pir:[I hilus i [DB:pi al trar bct2]	N:E640 nfluen r2] >g nsmembra DE:Hae	10] [Adzae con p:[GI:gane pro mophilo	C:E640 nserve g15735 otein] us inf	10] d hypoth 83] [LN: [GN:HI0 luenzae	etical U32741] 594] Rd section
ORF Name A17503000984_26774137_c2_574 Description NO-HIT	<u>NTID</u> 706	AAID 4478	NT LengthI	AA Length	score	probab	ility
ORF Name	NTID	AAID	<u>NT</u> LengthI	<u>AA</u> Length	score	probab	ility
AI7503000984_272593_c3_636	707	4479	879	292	82	0.0082	
Description sp:[LN:YORB_LISMO] [AC:P33382] [OR:L PROTEIN IN PLCB-LDH INTERGENIC REGIO >pir:[LN:143868] [AC:143868] [PN:OR monocytogenes] [DB:pir2] >gp:[GI:g14 monocytogenes] [SR:Listeria monocyto [DE:Listeria monocytogenes lecithina (plcB) gene complete cds, (ldh) gene [DI:complement]	N (ORFB FB] [CL 9648] [genes (se, lac)] [SP: :hypoth LN:LISA strain tate de	P33382] etical CTLDH] L028) I	DB:s prote: [AC:M8 [I] [I] [I]	swissp in MJ1 32881] DB:gen (actA)	rot] 413] [OR [OR:List pept-bct] gene com	:Listeria teria 1] plete cds,

ORF Name	NTID	AAID	NT AA score probability
A17503000984 2906307 f1 60	708	4480	
Description			JlJ
NO-HIT			
ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000984_2928437_c1_477	709	4481	921 306 375 1.4e-34
Description			JI
pir:[LN:A69401] [AC:A69401] [PN:con [OR:Archaeoglobus fulgidus] [DB:pir2 [AC:AE001021:AE000782] [PN:conserved [OR:Archaeoglobus fulgidus] [DB:genp 172 of the complete genome.] [NT:sim [LE:12088] [RE:13029] [DI:complement] >gp:[0 hypothe ept-bct2 ilar to	GI:g26 etical 2] [DE	49377] [LN:AE001021] protein] [GN:AF1210] :Archaeoglobus fulgidus section 86 of
ORF Name	NTID	AAID	NT AA score probability
A17503000984_2929718_c1_463	710	4482	1959 652 3345 0.0
Description			
<pre>gp:[GI:g3789932] [LN:AF090142] [AC:A [OR:Staphylococcus epidermidis] [DB: epidermidis lipase precursor (gehD) [DI:direct]</pre>	genpept-	bct2]	[EC:3.1.1.3] [DE:Staphylococcus
ORF Name	NTID	AAID	NT AA score probability
A17503000984_29398437_t2_132	711	4483	77
Description			
NO-HIT			
ORF Name	NTID	AAID	NT AA score probability
AI7503000984_3003137_f2_215	712	4484	
Description			
pir:[LN:H70313] [AC:H70313] [PN:cob. [OR:Aquifex aeolicus] [DB:pir2] >gp: [PN:cobalamin synthesis related prot. [DB:genpept-bct2] [DE:Aquifex aeolicus:[LE:11287] [RE:12165] [DI:direct]	[GI:g298 ein CobW	32874]] [GN	[LN:AE000675] [AC:AE000675:AE000657] :cobW] [OR:Aquifex aeolicus]
ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000984_30103592_c3_600	713	4485	132 43
Description			J
NO-HIT			

A17503000984_30251551_f3_412
sp:[LN:YO21_BPHP1] [AC:P51723] [OR:BACTERIOPHAGE HP1] [DE:HYPOTHETICAL 19.2 KD PROTEIN
IN REP-HOL INTERGENIC REGION (ORF21)] [SP:P51723] [DB:swissprot] >pir:[LN:S69527] [AC:S69527] [PN:hypothetical protein 21] [OR:phage HP1] [DB:pir2] >gp:[GI:g1046248] [LN:BHU24159] [AC:U24159:U06847:M28366:M12911:M22941:M12910:M15313] [OR:Bacteriophage HP1] [DB:genpept-phg] [DE:Bacteriophage HP1 strain HP1c1, complete genome.] [NT:orf21]
[LE:17028] [RE:17528] [DI:direct]
A17503000984_30271882_c3_618 715 4487 720 239 268 3.0e-23
Description gp:[GI:g490316] [LN:A02585] [AC:A02585] [GN:bioD] [OR:synthetic construct] [DB:genpept-pat] [DE:Synthetic (LORF1) bioD gene.] [LE:39] [RE:764] [DI:direct]
A17503000984_30272531_f2_228 716 4488 144 47 Description NO-HIT
ORF Name NTID AAID NT AA LengthLength score probability A17503000984 30351677 c3 598 717 4489 804 267
Description Page 1975
NO-HIT
AI7503000984_30742307_c2_572
Description pir:[LN:F69670] [AC:F69670] [PN:glycine betaine/carnitine/choline ABC transporter (membrane p) opuCD] [GN:opuCD] [CL:glycine betaine/carnitine/choline ABC transporter] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:g2271392] [LN:AF009352] [AC:AF009352] [PN:transmembrane protein] [GN:opuCD] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis osmoprotectant transport system OpuC includingATPase (opuCA), transmembrane protein (opuCB), osmoprotectantbinding protein precursor (opuCC) and transmembrane protein (opuCD)genes, complete cds.] [NT:OpuCD; part of the osmoprotectant transport system] [LE:3627] [RE:4316] [DI:direct] >gp:[GI:e1186068:g2635893] [LN:BSUB0018] [AC:299121:AL009126] [PN:glycine betaine/carnitine/choline ABC] [GN:opuCD] [FN:high affinity transport of glycine betaine,] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 18 of 21): from 3399551to 3609060.] [NT:alternate gene name: yvbB] [LE:67059] [RE:67748] [DI:complement]
ORF Name NTID AAID NT AA score probability
A17503000984_3125687_f1_114
Description NO-HIT

ORF Name	NTID	AAID	NT AA LengthLength score probability
AI7503000984_31287513_c1_464	720	4492	150 49
Description			
NO-HIT	SAAA		
ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000984_3163552_c3_597	721	4493	963 320 356 1.4e-32
Description			
- ·	ccus aur tive sub e domain p-1C), c ansporte	reus] ostrat oligop erputa	[DB:genpept-bct2] [DE:Staphylococcus ebinding domain (opp-1A), oligopeptide -1B), oligopeptide transporter eptide transporterputative ATPase tive ATPase domain (opp-1F) genes,
ORF Name	NTID	AAID	NT AA score probability
A17503000984_3174187_f2_222	722	4494	Length Length [109]
Description		1171	
NO-HIT			
			ATTO 2.2
ORF Name	NTID	AAID	NT AA Score probability
A17503000984_32609682_f3_403	723	4495	129 42
Description	<u> </u>		
NO-HIT			
ORF Name	NTID	AAID	NT AA LengthLength
A17503000984_32615677_c2_544	724	4496	1035 344 510 6.7e-49
Description		•	
[RE:22380] [DI:direct] >gp:[GI:e1182	>gp:[GI: R:Bacill bct1] [D ologue o 237:g263 subtilis rom 1946	d1023 us su E:Bac f adh 2571]] [DB 51 to	108:g2415736] [LN:AB000617] btilis] [SR:Bacillus subtilis illus subtilis genomic DNA, 22 to 25 esion protein precursor of] [LE:21421] [LN:BSUB0002] [AC:Z99105:AL009126] :genpept-bct1] [DE:Bacillus subtilis
ORF Name AI7503000984 32755 12 185	<u>NTID</u>	<u>AAID</u> 4497	NT AA score probability LengthLength 54
Description	[123]	147/	
NO-HIT			
ORF Name	NTID	AAID	NT AA score probability
A17503000984_33241562_£2_147	726	4498	144 47
Description			
NO-HIT			

			2777	7. 7.	
ORF Name	NTID	AAID	NT LengthL	ength score	probability
A17503000984_33479716_c3_634	727	4499	7 1401 4	66 421	1.8e-39
Description	L		JL		<u> </u>
gp:[GI:e1429016:g4753872] [LN:SCH10] protein] [GN:SCH10.26c] [OR:Streptom coelicolor cosmid H10.] [NT:SCH10.26 [LE:22729] [RE:24183] [DI:complement	yces co	elicol	or] [DB:	genpept-bct	1] [DE:Streptomyces
ORF Name	NTID	AAID	<u>NT</u> LengthL	AA ength score	probability
A17503000984_34171927_c1_468	728	4500	144 4	7	
Description	L				
NO-HIT					
ORF Name	NTID	AAID		AA ength	probability
A17503000984_34277062_c3_602	729	4501	153 5	0	
<u>Description</u>					
NO-HIT					
ORF Name	NTID	AAID	NT LengthL	AA ength score	probability
A17503000984_34408552_f1_21	730	4502	126 4	1	
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> LengthL	AA ength score	probability
AI7503000984_34429837_t1_39	731	4503	948 3	15 769	2.4e-76
Description sp:[LN:ARCL_ECOLI] [AC:Q46807] [GN:Y KINASE-LIKE PROTEIN 1] [SP:Q46807] [[PN:hypothetical protein b2874] [CL: >gp:[GI:g887824] [LN:ECU28375] [AC:U [DE:Escherichia coli K-12 genome; ap [LE:21276] [RE:22208] [DI:direct] >g [PN:putative kinase] [GN:yqeA] [FN:p coli] [DB:genpept-bct2] [DE:Escheric completegenome.] [NT:o310; This 310 [RE:12292] [DI:direct]	DB:swiss carbamat 28375] proximat p:[GI:gi utative hia col:	sprot] te kin [OR:Es tely 6 178923 enzym	>pir:[Lase] [OR cherichi 4 to 65 8] [LN:Ae; Not community MG1655 pct iden	N:B65071] [:Escherichi a coli] [DB minutes.] [E000370] [A lassified] section 260 tical (21 g	AC:B65071] a coli] [DB:pir2] :genpept-bct1] NT:ORF_0310] C:AE000370:U00096] [OR:Escherichia of 400 of the
ORF Name	NTID	AAID	<u>NT</u> <u>Length</u> Le	AA ength	probability
A17503000984_34631527_£1_109	732	4504	336 1	11 374	1.7e-34
Description					
<pre>gp:[GI:d1045996:g5360820] [LN:D86934 aureus] [SR:Staphylococcus aureus (s [DB:genpept] [DE:Staphylococcus aure</pre>	train:N3	315) DI	NA, clon	e_lib:libra	ry of N31]

[DE:Staphylococcus aureus genes, mec region, [NT:ORF N026; putative] [LE:19527] [RE:19751] [DI:direct]

ORF Name	NTID	AAID NT AA score probability				
A17503000984_34642135_c3_663	733	4505 246 81 79 0.020				
Description gp:[GI:g488889] [LN:A12521] [AC:A12521] [PN:Acidic Basic Repeat Antigen Rhoptry (ABRA)] [OR:Plasmodium falciparum] [SR:malaria parasite P. falciparum] [DB:genpept-pat] [DE:Ag189 clone.] [LE:1:61] [RE:45:963] [DI:directJoin]						
	pt-bct2] [DE:Staphylococcus aureus surface protein lin resistant; contains a DS repeat area]				
ORF Name AI7503000984_35312766_f2_151 Description NO-HIT	NTID 735	AAID LengthLength score probability 4507 195 64				
[OR:Drosophila melanogaster] [SR:fru	it fly]	AAID NT AA LengthLength score probability 4508 318 105 81 0.012 [PN:BcDNA.GH10614] [GN:BcDNA.GH10614] [DB:genpept-inv2] [DE:Drosophila cDNA.GH10614) mRNA, complete cds.] [LE:14]				
	pt-bct2	AAID NT AA score probability 4509 3246 1081 2306 1.8e-251 [PN:surface protein Pls] [GN:pls] [DE:Staphylococcus aureus surface protein lin resistant; contains a DS repeat area]				

NT ORF Name NTID AAID probability score LengthLength AI7503000984 36135752 £1 90 738 4510 1128 Description sp:[LN:YPDA BACSU] [AC:P50736] [GN:YPDA] [OR:BACILLUS SUBTILIS] [DE:HYPOTHETICAL 36.3 KD PROTEIN IN RECQ-CMK INTERGENIC REGION] [SP:P50736] [DB:swissprot] >pir:[LN:A69934] [AC:A69934] [PN:thioredoxin reductase homolog ypdA] [GN:ypdA] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1183740:g2634713] [LN:BSUB0012] [AC:Z99115:AL009126] [GN:ypdA] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 12 of 21): from 2195541to 2409220.] [NT:similar to thioredoxin reductase] [SP:P50736] [LE:204681] [RE:205655] [DI:complement] >qp:[GI:e1185564:q2634730] [LN:BSUB0013] [AC:Z99116:AL009126] [GN:ypdA] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 13 of 21): from 2395261to 2613730.] [NT:similar to thioredoxin reductase] [SP:P50736] [LE:4961] [RE:5935] [DI:complement] >gp:[GI:g1146207] [LN:BACSERA] [AC:L47648] [GN:ypdA] [OR:Bacillus subtilis] [DB:genpept-bct2] [DE:Bacillus subtilis phosphoglycerate dehydrogenase (serA), ypaA, ferredoxin (fer), ypbB, recS, ypbD, ypbE, ypbF, ypbG, ypbH,glutamate dehydrogenase (ypcA), ypdA, ypdB, ypdC, spore cortexlytic enzyme (sleB), ypeB, ypfA, ypfB, cytidine monophosphatekinase (cmk), ypfD, ypgA, yphA, yphB, yphC, NAD+ dependentglycerol-3-phosphate dehydrogenase (glyc), yphE and yphF genes, complete cds.] [NT:putative] [LE:10742] [RE:11716] [DI:direct] NT AA AAID ORF Name NTID score probability LengthLength AI7503000984_36601703_c1_467 739 4511 939 312 7.<mark>0e-38</mark> 406 Description sp:[LN:APBE TREPA] [AC:083774] [GN:APBE:TP0796] [OR:TREPONEMA PALLIDUM] [DE:THIAMINE BIOSYNTHESIS LIPOPROTEIN APBE PRECURSOR] [SP:083774] [DB:swissprot] >pir:[LN:C71281] [AC:C71281] [PN:conserved hypothetical protein TP0796] [GN:TP0796] [CL:hypothetical protein HI0172] [OR:Treponema pallidum subsp. pallidum] [SR:, syphilis spirochete] [DB:pir2] >gp:[GI:g3323101] [LN:AE001250] [AC:AE001250:AE000520] [PN:conserved hypothetical protein] [GN:TP0796] [OR:Treponema pallidum] [DB:genpept-bct2] [DE:Treponema pallidum section 66 of 87 of the complete genome.] [NT:similar to GB:L42023 SP:P44550 PID:1003244] [LE:10082] [RE:11170] [DI:complement]

 ORF Name
 NTID
 AAID
 NT AA Length Length
 Score
 probability

 A17503000984_3939215_f2_247
 740
 4512
 984
 327
 377
 8.3e-35

 Description

gp:[GI:g2766193] [LN:SHU75349] [AC:U75349] [PN:periplasmic-iron-binding protein BhiC]
[GN:bhi operon] [OR:Brachyspira hyodysenteriae] [DB:genpept-bct2] [DE:Serpulina
hyodysenteriae bhi operon, complete sequence.] [LE:1674] [RE:2693] [DI:direct]

ORF Name	NTID	AAID NT AA score probability
A17503000984_3948408_c3_655	741	4513 777 258 315 3.1e-28
>gp:[GI:g144297] [LN:CDCXYNAB] [AC:M [SR:C.saccharolyticum DNA, clone pNZ xylanase A (XynA), beta-xylosidase (cds.] [NT:acetyl esterase (XynC)] [LN:AF005383] [AC:AF005383] [PN:acet saccharolyticus] [DB:genpept-bct2] [transport protein(XynG), putative tr	TICUS] swisspr ocellum [34459] 1400] [XynB) a E:1257] ylxylos DE:Cald ansport ynA), a	[EC:3.1] [DE:ACETYL ESTERASE, rot] >pir:[LN:B37202] [AC:B37202] m saccharolyticum] [EC:3.1.1.6] [DB:pir2] [OR:Caldicellulosiruptor saccharolyticus] [DB:genpept-bct1] [DE:C.saccharolyticum andacetyl esterase (XynC) genes, complete] [RE:2057] [DI:direct] >gp:[GI:g2645420] sidase] [GN:XynC] [OR:Caldicellulosiruptor dicellulosiruptor saccharolyticus putative t protein (XynH), xylanase (XynF),xylanase acetylxylosidase(XynC) and xylanase (XynB)
ORF Name	NTID	AAID NT AA score probability
<pre>subsp. cremoris] [DB:genpept-bct2] [insertion sequenceIS982 putative tra cluster(epsRXABCDEFGHIJKL), complete >gp:[GI:g2072447] [LN:AF036485] [AC:</pre>	DE:Lact nsposas cds.] AF03648 enpept]	
[GN:yvfR] [CL:ATP-binding cassette >gp:[GI:e1186097:g2635922] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-b 18 of 21): from 3399551to 3609060.] protein)] [LE:97865] [RE:98770] [DI: [AC:Z94043] [PN:hypothetical protein	homolog 018] [A ct1] [D [NT:sim complem] [GN:y omic DN	NA fragment (88 kb).] [NT:probable ABC ect]
ORF Name A17503000984_4079382_f3_313 Description NO-HIT	NTID 744	AAID NT AA score probability 4516 243 80
ORF Name A17503000984_4094052_f3_330 Description NO-HIT	<u>NTID</u> 745	AAID NT AA score probability 4517 219 72

ORF Name AI7503000984_4098385_f1_115 Description NO-HIT	NTID	<u>AAID</u> 4518	NT AA LengthLength 129 42
ORF Name AI7503000984_4101063_f1_113 Description pir: [LN:T02638] [AC:T02638] [PN:G2 discoideum] [DB:pir2] >gp:[GI:g30685 [GN:g2] [OR:Dictyostelium discoideum	87] [LN:] [DB:ge	homolo AF0005	580] [AC:AF000580] [PN:G2-like] -invl] [DE:Dictyostelium discoideum asmid Ddp1 g2/g3/d4 protein; possible]
ORF Name A17503000984_4101640_c2_545 Description sp:[LN:YQIG_BACSU] [AC:P54524] [GN:Y [DE:PROBABLE NADH-DEPENDENT FLAVIN O >pir:[LN:C69961] [AC:C69961] [PN:NA [GN:yqiG] [OR:Bacillus subtilis] [D [AC:D84432:D82370] [PN:YqiG] [OR:Bac (strain:JH642(trpC2 PheA1)) DNA] [DB region containing skin element.] [LE >gp:[GI:e1185689:g2634855] [LN:BSUBO [OR:Bacillus subtilis] [DB:genpept-b 13 of 21): from 2395261to 2613730.] oxidoreductase] [SP:P54524] [LE:1204	XIDOREDU DH-deper B:pir2] illus su :genpept :202096] 013] [AC ct1] [DE	R:BACII JCTASE ident f >gp:[0 ibtilis -bct1] [RE:2 C:Z9911 E:Bacil ilar to	YQIG,] [SP:P54524] [DB:swissprot] flavin oxidoreductase homolog yqiG] GI:d1013261:g1303926] [LN:BACJH642] g] [SR:Bacillus subtilis [DE:Bacillus subtilis DNA, 283 Kb 203214] [DI:complement] 16:AL009126] [GN:yqiG] [FN:unknown] flus subtilis complete genome (section on NADH-dependent flavin
[PN:8-amino-7-oxononanoate synthase [DB:genpept-bct2] [DE:Haemophilus in	2] [DB:shomolog] homolog]] >gp:[G (bioF)] fluenzae	4521 553] [C swisspr [CL:5 GI:g157 [GN:HI c Rd se	rot] >pir:[LN:D64129] [AC:D64129] 5-aminolevulinate synthase] 74397] [LN:U32830] [AC:U32830:L42023] [1553] [OR:Haemophilus influenzae Rd]
ORF Name A17503000984_4722131_f2_268 Description NO-HIT	NTID 750	<u>AAID</u> 4522	NT AA LengthLength 141 46

ORF Name	NTID AAID NT AA score probability
A17503000984_4765_f2_245	751 4523 321 106 90 0.0025
Description	
[DE:Tetrahymena pyriformis mitochond	AF160864] [PN:haem lyase] [GN:yejR] ormis] [SR:Tetrahymena pyriformis] [DB:genpept] drial DNA, complete genome.] [NT:ATA initiation [LE:18212] [RE:19750] [DI:complement]
ORF Name	NTID AAID NT AA score probability
A17503000984_4773518_c3_664	752 4524 885 294 563 1.6e-54
[FN:cell membrane lipoprotein] [OR:S	PP] [AC:Y12602] [PN:acid phosphatase] [GN:lppC] Streptococcus equisimilis] [DB:genpept-bct1] and lppC genes.] [LE:1390] [RE:2247] [DI:direct]
ORF Name	$rac{ ext{NTID}}{ ext{NTD}} = rac{ ext{AAID}}{ ext{LengthLength}} rac{ ext{AA}}{ ext{score}} = rac{ ext{probability}}{ ext{probability}}$
AI7503000984_4782963_c1_460	753 4525 1365 454 1105 6.0e-112
Description	
>gp:[GI:g2982887] [LN:AE000676] [AC:	nase] [OR:Aquifex aeolicus] [DB:pir2] :AE000676:AE000657] [PN:DAPA aminotransferase] :genpept-bct2] [DE:Aquifex aeolicus section 8 of 109
ORF Name	NTID AAID NT AA score probability
A17503000984_4797177_c2_566	754 4526 588 195 308 1.7e-27
Description	
ORF Name	NTID AAID NT AA score probability
A17503000984_4798202_c3_623	755 4527 816 271 103 0.023
Description	
[OR:Plasmodium falciparum] [DB:pir2] [AC:AE001420:AE001362] [PN:hypothetifalciparum] [SR:malaria parasite P.	ical protein] [GN:PFB0850c] [OR:Plasmodium falciparum] [DB:genpept-inv2] [DE:Plasmodium of 73 of thecomplete sequence.] [NT:predicted by
ORF Name	NTID AAID NT AA score probability
A17503000984_4800077_£3_350	756 4528 156 51
Description	

ORF Name A17503000984 4859628 c3 603	<u>NTID</u>	<u>AAID</u>	NT AA LengthLength score	probability
Description NO-HIT	[37]	1323		
ORF Name	NTID	AAID	NT AA LengthLength score	probability
A17503000984_4867842_c3_652	758	4530	978 325 486	2.3e-46
Description pir: [LN:E69400] [AC:E69400] [PN:3-h; [OR:Archaeoglobus fulgidus] [DB:pir2] [AC:AE001021:AE000782] [PN:3-hydroxy; [OR:Archaeoglobus fulgidus] [DB:genperson of the complete genome.] [NT:sim:[LE:9126] [RE:10073] [DI:complement]] >gp:[0 acyl-Co <i>l</i> ept-bct2	GI:g26 A dehy 2] [DE	49379] [LN:AE001021 drogenase (hbd-8)] :Archaeoglobus fulc	l] [GN:AF1206] gidus section 86 of
ORF Name	NTID	AAID	NT AA score	probability
AI7503000984_4884812_f3_314	759	4531	1257 418 579	1.1e-78
Description				
<pre>gp:[GI:e1358508:g3980137] [LN:LMO3463 desuccinylase] [GN:dapE] [OR:Listeria monocytogenes ascB, inlG, inlH, inlE,</pre>	a monocy	togen/	es] [DB:genpept-bct	[DE:Listeria
ORF Name	NTID	AAID	NT AA LengthLength score	probability
A17503000984_4895061_c2_561	760	4532	2271 756 2671	6.8e-278
Description pir:[LN:S01788] [AC:S01788:B32305:F64 formate-lyase I] [GN:pflB:pfl] [CL:fl homology] [OR:Escherichia coli] [EC:2 >gp:[GI:d1036624:g1651427] [LN:D90728 c-acetyltransferase (EC 2.3.1.54).] coli(strain:K12) DNA, clone:Kohara cl genomic DNA. (20.4 - 20.8 min).] [NT: [LE:6965] [RE:9247] [DI:complement] > [OR:Escherichia coli] [DB:genpept-bct (EC 2.3.1.54).] [NT:pyruvate formate- [DI:direct] >gp:[GI:g1787131] [LN:AEC acetyltransferase 1] [GN:pflB] [FN:er [OR:Escherichia coli] [DB:genpept-bct section 82 of 400 of the completegence P09373] [LE:5588] [RE:7870] [DI:comple	Formate 2.3.1.54 3] [AC:E [GN:pf1] Lone #21 :ORF_ID: pgp:[GI: -lyase (000192] nzyme; E 2] [EC: ome.] [N	C-ace [DB 90728 [OR:] 6] [DI 0216# g42370 E. co. [AA 1- [AC:AI cnergy 2.3.1	tyltransferase 1:gl :pir1] [MP:20.5] :AB001340] [PN:Form Escherichia coli] [B:genpept-bct1] [DE 7; similar to PIR A D] [LN:ECPFL] [AC:X Li pfl gene for pyr 760)] [SP:P09373] [E000192:U00096] [PN metabolism, carbon .54] [DE:Escherichi	Eycyl radical Thate [SR:Escherichia E:Escherichia coli Eccession Number] [08035] Euvate formate-lyase [LE:101] [RE:2383] E:formate E: Anaerobic] Ea coli K-12 MG1655
ORF Name	NTID	AAID	NT AA LengthLength	probability
AI7503000984_4974091_c2_569	761	4533		0.00087
Description pir:[LN:B71359] [AC:B71359] [PN:cons [OR:Treponema pallidum subsp. pallidu >gp:[GI:g3322423] [LN:AE001200] [AC:A protein] [GN:TP0156] [OR:Treponema pa section 16 of 87 of the complete genc PID:1003656] [LE:2984] [RE:3388] [DI:	um] [SR: AE001200 illidum] ome.] [N	, sypl :AE000: DB:g T:simi	nilis spirochete] [0520] [PN:conserved genpept-bct2] [DE:T	DB:pir2] hypothetical reponema pallidum

ORF Name	NTID	AAID LengthLength score probability	<u>′</u>				
A17503000984_505301_f3_345	762	4534 291 96					
Description							
NO-HIT							
	The second secon	NTT 7.7.					
ORF Name	NTID	AAID LengthLength score probability	_				
A17503000984_5109785_c3_670	763	4535 255 84 76 0.010					
Description gp:[GI:g4103231] [LN:AF021085] [AC:AF021085] [PN:cytochrome b] [GN:cytb] [OR:Mitochondrion Edaphus sp.] [SR:Edaphus sp] [DB:genpept-inv2] [DE:Edaphus sp. cytochrome b (cytb) gene, mitochondrial gene encodingmitochondrial protein, partial cds.] [LE:<1] [RE:>465] [DI:direct]							
ORF Name	NTID	$rac{ ext{AAID}}{ ext{LengthLength}} rac{ ext{SCORe}}{ ext{probability}}$					
A17503000984_5113413_c3_653	764	4536 642 213 579 3.3e-56					
Description							
pir:[LN:D69670] [AC:D69670] [PN:glycine betaine/carnitine/choline ABC transporter (membrane p) opuCB] [GN:opuCB] [CL:glycine betaine/carnitine/choline ABC transporter] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:g2271390] [LN:AF009352] [AC:AF009352] [PN:transmembrane protein] [GN:opuCB] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis osmoprotectant transport system OpuC includingATPase (opuCA), transmembrane protein (opuCB), osmoprotectantbinding protein precursor (opuCC) and transmembrane protein (opuCD)genes, complete cds.] [NT:OpuCB; part of the osmoprotectant transport system] [LE:2025] [RE:2678] [DI:direct] >gp:[GI:e1186070:g2635895] [LN:BSUB0018] [AC:Z99121:AL009126] [PN:glycine betaine/carnitine/choline ABC] [GN:opuCB] [FN:high affinity transport of glycine betaine,] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 18 of 21): from 3399551to 3609060.] [NT:alternate gene name: yvbD] [LE:68697] [RE:69350] [DI:complement]							
ORF Name	NTID	$rac{ ext{NT}}{ ext{AAID}} rac{ ext{NT}}{ ext{Length}} rac{ ext{score}}{ ext{tore}}$					
A17503000984_5160925_c1_443	765	4537 [192] [63					
Description NO-HIT							
ORF Name	NTID	AAID <u>NT AA</u> LengthLength score probability					
AI7503000984_5189037_c3_620	766	4538 711 236 351 4.7e-32					
Description							
[DB:pir2] [MP:FOR1244632-1245345] > [PN:6-carboxyhexanoate-CoA ligase (b [DB:genpept-bct2] [DE:Methanococcus	ioW] [OF gp:[GI:g ioW)] [G jannasch	xanoateCoA ligase,] :Methanococcus jannaschii] [EC:6.2.1. 1591935] [LN:U67570] [AC:U67570:L7711 N:MJ1297] [OR:Methanococcus jannaschi ii section 112 of 150 of the complete PID:142594 percent] [LE:6098] [RE:681	7] i]				
ORF Name	NTID	AAID <u>NT AA</u> LengthLength score probability					
AI7503000984_5860630_c2_564	767	4539 171 56					
Description NO-HIT							

ORF Name	NTID AAII	NT AA LengthLength score	probability
AI7503000984 6056567_c1_490	768 4540		9.1e-29
Description	L L		
pir: [LN:D64866] [AC:D64866] [PN:hyp [DB:pir2] >gp: [GI:g1787449] [LN:AE00 dihydroxyacetone kinase (EC 2.7.1.2) [OR:Escherichia coli] [DB:genpept-bc 400 of the completegenome.] [NT:f210 [LE:4528] [RE:5160] [DI:complement]	0218] [AC:AE] [GN:b1199] t2] [DE:Esch	000218:U00096] [PN:p [FN:putative enzyme erichia coli K-12 MG	utative ; Not classified] 1655 section 108 of
ORF Name	NTID AAID	NT AA LengthLength	probability
AI7503000984_625262_c1_430	769 4541	. 228 75 54	0.013
Description			
gp:[GI:g765037] [LN:DROMTTRND] [AC:M [OR:Mitochondrion Drosophila melanog [DE:D.melanogaster Trp-tRNA, Cys-tRN cytochrome oxidase subunit 1 (5' end	aster] [SR:f A, Tyr-tRNA,	ruit fly] [DB:genpep NADH dehydrogenases	t-inv2] ubunit 2 (3' end)
ORF Name	NTID AAID	NT AA LengthLength	probability
A17503000984_6302217_c3_638	770 4542	948 315 1335	2.5e-136
Description			
membrane] [GN:opp-1B] [OR:Staphyloco aureus oligopeptide transporter puta transporter putative membranepermeas putativemembrane permease domain (op domain (opp-1D), and oligopeptide tr complete cds; and unknowngene.] [LE:	tive substra e domain (op p-1C), oligo ansporterput	tebinding domain (op p-1B), oligopeptide peptide transporterp ative ATPase domain	p-1A), oligopeptide transporter utative ATPase
ORF Name	NTID AAID	NT AA LengthLength	probability
A17503000984_6725817_c3_658	771 4543	165 54	
Description			
NO-HIT			
ORF Name	NTID AAID	NT AA LengthLength	probability
A17503000984_7072825_c1_435	772 4544	774 257 459	1.7e-43
Description			
gp:[GI:g4980796] [LN:AE001711] [AC:A [GN:TM0297] [OR:Thermotoga maritima] 23 of 136 of the complete genome.] [identity:] [LE:7161] [RE:7934] [DI:d	[DB:genpept NT:similar t	-bct2] [DE:Thermotoga	a maritima section
ORF Name	NTID AAID	<u>NT AA</u> LengthLength	probability
A17503000984_800300_c1_458	773 4545		5.0e-28
Description	<u> </u>		
	_	brane-associated ant:	igen (clone pPf

ORF Name NTID AAID NT AA score probability
A17503000984_814140_c1_437
Description Sp:[LN:UHPT_ECOLI] [AC:P13408:P76727] [GN:UHPT] [OR:ESCHERICHIA COLI] [DE:HEXOSE PHOSPHATE TRANSPORT PROTEIN] [SP:P13408:P76727] [DB:swissprot] >pir:[LN:MMECHP] [AC:A30395:H41853:C65168:Q00500:S30079] [PN:hexose phosphate transport protein uhpT] [GN:uhpT] [CL:hexose phosphate transport protein uhpT] [OR:Escherichia coli] [DB:pir1] [MP:82 min] >gp:[GI:g148115] [LN:ECOUHP] [AC:M17102] [OR:Escherichia coli] [SR:E.coli DNA, clone pRJK10] [DB:genpept-bct1] [DE:E.coli uhp operon encoding UhpA, UhpB, UhpC, and UhpT protein, (encoding hexose phosphate transport protein), complete cds, and anilvBN operon encoded protein, 3' end.] [NT:hexose phosphate transport protein UhpT] [LE:3722] [RE:5113] [DI:direct] >gp:[GI:g148120] [LN:ECOUHPABCT] [AC:M89479] [PN:hexosephosphate transport protein] [GN:uhpT] [OR:Escherichia coli] [SR:Escherichia coli DNA] [DB:genpept-bct1] [DE:Escherichia coli uhpABCT operon encoding nexosephosphateutilization protein (uhpA) gene, complete cds, and nexosephosphatetransport protein (uhpB, uhpC, uhpT) genes, complete cds.] [LE:3722] [RE:5113] [DI:direct] >gp:[GI:g2367259] [LN:AE000444] [AC:AE000444:U00096] [PN:hexose phosphate transport protein] [GN:uhpT] [FN:transport; Transport of small molecules:] [OR:Escherichia coli] [DB:genpept-bct2] [DE:Escherichia coli K-12 MG1655 section 334 of 400 of the completegenome.] [NT:f463; 99 pct identical amino acid sequence and] [LE:5590] [RE:6981] [DI:complement]
NTID AAID NT AA LengthLength Score probability A17503000984_81525_f3_398 Pescription NO-HIT
NTID AAID NT AA Score probability AI7503000984_816878_c3_637 Pescription Gescription Gescrip
NTID AAID NT AA LengthLength score probability AI7503000984_822150_f2_262 777
NTID AAID NT AA score probability A17503000984_875765_c2_525

ORF Name	NTID	AAID	NT AA LengthLength score probability
AI7503000984_969157_c1_420	779	4551	1026 341 220 8.5e-16
	occus a tive su e domai p-1C), ansport	ureus] bstrat n (opp oligop erputa	[DB:genpept-bct2] [DE:Staphylococcus ebinding domain (opp-1A), oligopeptide -1B), oligopeptide transporter eptide transporterputative ATPase tive ATPase domain (opp-1F) genes,
ORF Name	NTID	AAID	NT AA score probability
AI7503000984_9806718_c1_474	780	4552	762 253 862 3.4e-86
	ylococo de tran utative brane p pp-1D),	us aur sporte membr ermeas and o	eus] [DB:genpept-bct2] r putative substratebinding domain anepermease domain (opp-1B),
ORF Name	NTID	AAID	NT AA score probability
AI7503000984_984686_c1_478	781	4553	
Description pir:[LN:C69670] [AC:C69670] [PN:gly((ATP-bindin) opuCA] [GN:opuCA] [CL:(ATP-binding cassette homology:CBS homology:[GI:g2271389] [LN:AF009352] [AC:(subtilis] [DB:genpept-bct1] [DE:BacilincludingATPase (opuCA), transmembrane precursor (opuCC) and transmembrane part of the osmoprotectant transport part of the osmoprotectant part of th	glycine mology] AF00935: llus sui ne prote protein system 018] [A6 puCA] [3: genpep	betair [OR:B: 2] [PN btilis ein (opuCl] [LE:: C:Z991: FN:high	ne/proline transport protein proV: acillus subtilis] [DB:pir2] :ATPase] [GN:opuCA] [OR:Bacillus osmoprotectant transport system OpuC puCB), osmoprotectantbinding protein D)genes, complete cds.] [NT:OpuCA; B60] [RE:2002] [DI:direct] 21:AL009126] [PN:glycine n affinity transport of glycine [DE:Bacillus subtilis complete
ORF Name	NTID	AAID	NT AA score probability
AI7503000984_9970167_f3_393	782	4554	861 286 219 2.9e-25
[GN:yesN] [CL:response regulator hor >gp:[GI:e1182675:g2633009] [LN:BSUB00	mology] 004] [Ad ctl] [D] :simila:	OR:Ba 2:Z9910 E:Baci	07:AL009126] [GN:yesN] [FN:unknown] Llus subtilis complete genome (section wo-component response regulator
ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000984_9973515_c1_495	783	4555	261 86
Description NO-HIT			

ORF Name	NTID	AAID LengthLength score probability
A17503000985_10000183_c3_2061	784	4556 168 55
Description		<u>, </u>
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000985_10034627_f3_956	785	4557 144 47
Description		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
AI7503000985_10195252_£2_807	786	4558 126 41
Description		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000985_10203501_c2_1757	787	4559 156 51
Description		
NO-HIT .		
ORF Name	NTID	AAID NT AA score probability LengthLength
A17503000985_10241433_c1_1640	788	4560 126 41 104 1.4e-05
[AC:C69783] [PN:NADH dehydrogenase [OR:Bacillus subtilis] [DB:pir2] >gp [GN:ydgI] [OR:Bacillus subtilis] [SR [DB:genpept-bct1] [DE:Bacillus subtiregionbetween 35 and 47 degree.] [NT [RE:146039] [DI:complement] >gp:[GI:[AC:Z99107:AL009126] [GN:ydgI] [FN:ult] [DE:Bacillus subtilis complete genomes.]	YDGI,] homolog :[GI:d1 ::Bacill dis gen ::SIMILA e118254 unknown] ne (sect	[SP:P96707] [DB:swissprot] >pir:[LN:C69783] g ydgI] [GN:ydgI] [CL:nitroreductase] 1020152:g1881372] [LN:AB001488] [AC:AB001488] lus subtilis (strain:168) DNA] nome sequence, 148 kb sequence of the AR TO NITROREDUCTASE.] [LE:145410]
ORF Name	NTID	AAID <u>NT AA</u> score probability <u>LengthLength</u>
A17503000985_10332262_t1_254	789	4561 366 121
Description NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000985_1046885_f1_82	790	4562 2217 738 1356 1.5e-138
Description		
-	ept-bct gA, ORF	

ORF Name	NTID	AAID NT AA score probability
A17503000985_10585432_c1_1539	791	4563 591 196 171 2.3e-18
[OR:Bacillus subtilis] [EC:2.3.2.2] [AC:Z99113:AL009126] [PN:gamma-gluta metabolism] [OR:Bacillus subtilis] [complete genome (section 10 of 21): pac] [SP:P54422] [LE:222733] [RE:224 [LN:BSUB0011] [AC:Z99114:AL009126] [FN:glutathione metabolism] [OR:Baci [DE:Bacillus subtilis complete genom [NT:alternate gene name: pac] [SP:P5 >gp:[GI:g1491813] [LN:BSU49358] [AC: [OR:Bacillus subtilis] [SR:Bacillus	T] [OR:I CURSOR,] C4504] ase] [GI [DB:pir2 myltrans DB:genpe from 178 496] [DI PN:gamma llus sub e (sect: 4422] [I U49358] subtilis	BACILLUS SUBTILIS] [EC:2.3.2.2] [SP:P54422] [DB:swissprot] [PN:gamma-glutamyltransferase, N:ggt] [CL:gamma-glutamyltransferase] [PS:P54422] [DB:swissprot] [PN:gamma-glutamyltransferase] [Sp:[GI:e1183499:g2634224] [LN:BSUB0010] [Speptidase] [GN:ggt] [FN:glutathione [Speptidase] [GN:ggt] [DE:Bacillus subtilis [S1201to 2014980.] [NT:alternate gene name: [I:direct] >gp:[GI:e1185314:g2634235] [A-glutamyltranspeptidase] [GN:ggt] [DB:genpept-bct1] [EC:2.3.2.2] [ion 11 of 21): from 2000171to 2207900.] [LE:3763] [RE:5526] [DI:direct] [PN:gamma-glutamyltranspeptidase] [GN:ggt]
ORF Name AI7503000985_10600010_c2_1931 Description NO-HIT	NTID 792	AAID NT AA score probability LengthLength 4564 192 63
ORF Name A17503000985_10601625_c3_2039 Description gp:[GI:e1286089:g3036830] [LN:CJAJ08 [GN:kpsM] [OR:Campylobacter jejuni] kpsT genes.] [LE:134] [RE:916] [DI:d	[DB:genp	-
[OR:Bacillus subtilis] [DB:genpept-b	ctl] [DI	AAID NT AA LengthLength score probability 4566 747 248 706 1.1e-69 :putative ATP binding subunit] [GN:ORF3] E:B.subtilis putative amino acid transporter [SP:P39456] [LE:1627] [RE:2370] [DI:direct]
ORF Name A17503000985_1063552_f3_982 Description NO-HIT	NTID 795	AAID NT AA score probability 4567 390 129
ORF Name A17503000985_1064050_f2_826 Description NO-HIT	NTID 796	AAID NT AA score probability 4568 150 49

ORF Name A17503000985_10657827_c1_1607	<u>NTID</u> 797	<u>AAID</u>	NT AA LengthLength score probability 1401 466 393 1.7e-36
Description gp:[GI:e304997:g2294506] [LN:A37836] [DB:genpept-pat] [DE:Sequence 6 from [LE:103] [RE:1689] [DI:direct]			
ORF Name	NTID	AAID	NT AA LengthLength score probability
AI7503000985_10667002_c2_1653 Description NO-HIT	798	4570	129 42
MO-HII			
ORF Name	NTID	AAID	NT AA LengthLength
A17503000985_10667003_f1_29	799	4571	174 57
Description NO-HIT			
ORF Name	NTID	AAID	NT AA LengthLength score probability
AI7503000985_10718762_f1_204	800	4572	165 54
Description		L	
NO-HIT			
ORF Name	NTID	AAID	NT AA LengthLength
A17503000985_10736312_c1_1452	801	4573	132 43
Description			
NO-HIT			
ORF Name	NTID	AAID	NT AA LengthLength
A17503000985_1074177_c2_1838	802	4574	825 274 417 4.8e-39
Description			
pir:[LN:F64819] [AC:F64819] [PN:hypo			
			[AC:D90719:AB001340] [PN:Hypothetical
<pre>protein 1] [OR:Escherichia coli] [SR clone #206] [DB:genpept-bct1] [DE:Esc</pre>			
[NT:ORF_ID:0207#5; similar to PIR Acc			
[DI:complement] >gp:[GI:d1036496:g406			
[PN:Hypothetical protein 1] [OR:Esche	erichia	coli]	[SR:Escherichia coli(strain:K12) DNA,
clone:Kohara clone #207] [DB:genpept-			
18.8 min).] [NT:ORF_ID:o207#5; similated [DI:complement] >gp:[GI:g1787043] [Li			
hypothetical protein] [GN:b0822] [FN:			
[DB:genpept-bct2] [DE:Escherichia col			
completegenome.] [NT:f271; This 271 a			
[RE:7424] [DI:complement]			

ORF Name	NTID AAID LengthLength score probability
A17503000985_10756925_±1_228	803 4575 1566 521 793 6.9e-79
[SR:Bacillus subtilis (strain:AC327)	8] [AC:D86418] [PN:YfnA] [OR:Bacillus subtilis]) DNA] [DB:genpept-bct1] [DE:Bacillus subtilis rtialsequence.] [LE:7539] [RE:8927] [DI:direct]
ORF Name	NTID AAID NT AA score probability
A17503000985_10938903_f1_322	804 4576 144 47
Description NO-HIT	
ORF Name	NTID AAID LengthLength score probability
A17503000985_10954127_£3_1025	805 4577 201 66
NO-HIT	
ORF Name	NTID AAID NT AA score probability
A17503000985_10973385_c3_2080	806 4578 312 103 80 0.010
[GN:DG1122] [OR:Dictyostelium discoi	AF111944] [PN:development protein DG1122] ideum] [DB:genpept-inv2] [DE:Dictyostelium DG1122 (DG1122)gene, partial cds.] [LE:207:829]
ORF Name	$rac{ ext{NTID}}{ ext{AAID}} = rac{ ext{NT}}{ ext{Length} ext{Length}} rac{ ext{score}}{ ext{probability}}$
AI7503000985_11063801_£1_381	807 4579 225 74
Description NO-HIT	
ORF Name	NTID AAID LengthLength score probability
A17503000985_11132010_c1_1519	808 4580 126 41
Description	
NO-HIT	
ORF Name	NTID AAID LengthLength score probability
A17503000985_115761_c3_2118	809 4581 1458 485 799 1.6e-79
[SR:PCC 6803, , PCC 6803] [SR:PCC 68 [LN:D90907] [AC:D90907:AB001339] [PN sp.] [SR:Synechocystis sp. (strain:Po	pothetical protein slr1363] [OR:Synechocystis sp.] 803,] [DB:pir2] >gp:[GI:d1018310:g1652657] N:hypothetical protein] [GN:glgP] [OR:Synechocystis PCC6803) DNA] [DB:genpept-bct1] [DE:Synechocystis 1056467-1188885.] [NT:ORF_ID:slr1363] [LE:49200]

ORF Name	NTID	AAID	NT Length	<u>AA</u> Lengtl	score	probability
A17503000985_117150_f1_173	810	4582	1311	436	1525	1.9e-156
Description			JLJ	L	الـــــا	
<pre>gp:[GI:g4096796] [LN:SCU40157] [AC:U [DB:genpept-bct2] [DE:Staphylococcus andSpoVE-like protein (orf2) genes, to] [LE:193] [RE:1362] [DI:direct]</pre>	carnos	us con	densing	g-enzyı	me-like	protein (orf1)
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Lengtl	score	probability
A17503000985_1173130_c1_1530	811	4583	2091	696	2666	2.3e-277
Description	<u></u>				الــــــا	
pir:[LN:S46952] [AC:S46952:S63605] glucose-specific, factor IIA:glucose phosphotransferase system enzyme II, II, glucose-specific, factor 1:prote glucose-specific, factor II] [GN:glc N-acetylglucosamine-specific enzyme enzyme II, factor II homology:phosph factor III homology] [OR:Staphylococ>gp:[GI:g1072418] [LN:SCGLCAB] [AC:X carnosus] [DB:genpept-bct1] [DE:S.ca [DI:direct]	permeas glucose in-Npi-p A:ptsG II:phosp otransfe cus carr 93360:X8	se:phoe-spec phosphosphological CL: photramerase nosus] 80415]	sphoend ific:phohistic phospho nsferas system [EC:2. [GN:gl	olpyrumosphot dines otransi se syst glucos (7.1.69	vate:gl transfe sugar p ferase tem glu se-spec 9] [DB: DR:Stap	ucose rase system enzyme chosphotransferase, system cose-specific cific enzyme II, pir1] chylococcus
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
AI7503000985_11756543_f2_497	812	4584	231	76]	
Description NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
AI7503000985_1178593_f3_990	813	4585	1590	529	1527	1.1e-156
Description					J I	
pir:[LN:C69794] [AC:C69794] [PN:glu] [OR:Bacillus subtilis] [DB:pir2] > [AC:Z99107:AL009126] [GN:yerD] [FN:u] [DE:Bacillus subtilis complete genom [NT:similar to glutamate synthase (f>gp:[GI:e1167974:g2577963] [LN:BSYER [OR:Bacillus subtilis] [DB:genpept-byerA to sapB gene.] [NT:similar to p [RE:4808] [DI:complement]	gp:[GI:enknown] e (sectierredoxi ABCD] [A	e11826; [OR:Baion 4 (in)] [1 AC:Y15; E:Baci	39:g263 acillus of 21): LE:1155 254] [F llus su	32973] s subti from [886] [F PN:YerI abtilis	[LN:BS ilis] [600701 RE:1171 D prote s 13kB	UB0004] DB:genpept-bct1] to813890.] 63] [DI:complement] in] [GN:yerD] DNA fragment, from
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000985_1180292_c1_1536	814	4586	1164	387	1050	4.0e-106
Description		<u> </u>				
pir:[LN:F70069] [AC:F70069] [PN:capare [GN:ywsC]] [OR:Bacillus subtilis] [DI [AC:Z99122:AL009126] [GN:ywsC] [FN:un [DE:Bacillus subtilis complete genome [NT:similar to capsular polyglutamate [DI:complement] >gp:[GI:e1184496:g26. [FN:unknown] [OR:Bacillus subtilis] (section 19 of 21): from 3597091to 38 biosynthesis] [LE:101413] [RE:102594]	B:pir2] nknown] e (secti e biosyr 36115] [DB:genr 809700.]	>gp:[0 [OR:Bation 19 nthesis [LN:BSU pept] [NT:s	GI:e118 acillus of 21) s] [LE: JB0019] [DE:Bac similar	4496:9 subti : from 101413 [AC:2	g263611 llis] [n 35970 B] [RE: Z99122: subtil	5] [LN:BSUB0019] DB:genpept-bct1] 91to 3809700.] 102594] AL009126] [GN:ywsC] is complete genome

ORF Name AI7503000985_11855463_c3_2075 Description pir: [LN:H69611] [AC:H69611] [PN:3'-cysH:3'-phosphoadenylylsulfate reduction reduc	ctase, the cillus sure of the cillus sure of the cillus sure of the cillus cill	hiored ubtili C:AJ00 illus ion.] C:Z991 ynthes s subt	ine 5'-1 oxin den s] [EC:1 0974] [1 subtilis [LE:548] 12:AL009 is] [OR ilis con	phosphore pendent 1.8.99 PN:puta s] [DB] [RE:: 9126] :Bacill	osulfat: PAF .4] [I ative :genpe 1249] [PN:ph lus su genon	PS reductase:PAPS DB:pir2] ept-bct1] [DI:direct] hosphoadenosine hbtilis]
ORF Name A17503000985_11881630_f2_658 Description NO-HIT	NTID 816	<u>AAID</u> 4588	NT LengthI	AA Length	score	probability
ORF Name AI7503000985_11886592_c2_1671 Description sp:[LN:HIS2_HAEIN] [AC:P44434] [GN:HER] [EC:3.5.4.19:3.6.1.31] [DE:PYROPHOSE > pir:[LN:A64071] [AC:A64071] [PN:Pher] phosphoribosyl-ATP pyrophosphatase, lenzyme homology:hisI protein homology [EC:3.5.4.19:3.6.1.31] [DB:pir2] > gr [PN:phosphoribosyl-AMP cyclohydrolast [DB:genpept-bct2] [DE:Haemophilus in genome.] [NT:similar to SP:P06989 GR [DI:direct]	PHOHYDROI nosphorik [CL:his gy] [OR:F p:[GI:g15 se /] [GN nfluenzae	LASE,] cosyl- sI bif Haemop 573454 N:HI04 e Rd s	475] [OI [SP:P44] AMP cyclunctions hilus in] [LN:U3] 75] [OR:	R:HAEMO 4434] lohydro al enzy nfluenz 32730] :Haemon	[DB:swolase, yme:hi zae] [AC:U philus 163 of	vissprot] / sI bifunctional 32730:L42023] s influenzae Rd] the complete
ORF Name AI7503000985_11955127_f1_420	NTID 818	<u>AAID</u> 4590	NT LengthI	AA Length	score	probability

Description NO-HIT

NT NTID AAID probability ORF Name score LengthLength AI7503000985 119633 f1 70 4591 1389 462 Description pir:[LN:B69680] [AC:B69680:I40510] [PN:para-nitrobenzyl esterase,] [GN:pnbA] [CL:cholinesterase homology] [OR:Bacillus subtilis] [EC:3.1.1.-] [DB:pir2] >qp:[GI:q1762126] [LN:BSU46134] [AC:U46134] [PN:intracellular esterase B] [GN:estB] [OR:Bacillus subtilis] [DB:genpept-bct1] [EC:3.1.1.1] [DE:Bacillus subtilis putative orf1 unknown protein, putativetranscriptional regulator (slr), and intracellular esterase B(estB) genes, complete cds.] [NT:EstB; esterase of the serine-hydrolase family] [LE:1035] [RE:2504] [DI:direct] >gp:[GI:e1186127:g2635952] [LN:BSUB0018] [AC:Z99121:AL009126] [PN:para-nitrobenzyl esterase (intracellular] [GN:pnbA] [OR:Bacillus subtilis] [DB:genpept-bct1] [EC:3.1.1.1] [DE:Bacillus subtilis complete qenome (section 18 of 21): from 3399551to 3609060.] [NT:alternate gene name: estB] [LE:130145] [RE:131614] [DI:direct] >gp:[GI:e238702:g1495277] [LN:BSYVEFGNS] [AC:Z71928] [PN:para-nitrobenzyl esterase] [GN:pnbA] [OR:Bacillus subtilis] [DB:qenpept-bct1] [DE:B.subtilis pnbA, sigL, yve[J,K,L,M,N,O,P,Q,R,S,T] andyvf[A,B,C,D,E,F,G,H] genes.] [LE:132] [RE:1601] [DI:complement] >qp:[GI:e313129:q1945688] [LN:BSZ94043] [AC:Z94043] [PN:para-nitrobenzyl esterase] [GN:pnbA] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:B.subtilis genomic DNA fragment (88 kb).] [LE:50057] [RE:51526] [DI:complement] NTAA <u>sco</u>re ORF Name NTID probability LengthLength AI7503000985 1207287 c2 1669 4592 621 206 820 365 1.6e-33 Description sp:[LN:HIS1 LACLA] [AC:Q02129] [GN:HISG] [OR:LACTOCOCCUS LACTIS] [SR:, SUBSPLACTIS: STREPTOCOCCUS LACTIS] [EC:2.4.2.17] [DE:ATP PHOSPHORIBOSYLTRANSFERASE, [SP:Q02129] [DB:swissprot] >pir:[LN:D45734] [AC:D45734] [PN:HisG] [OR:Lactococcus lactis subsp. lactis] [DB:pir2] >gp:[GI:g2565141]

sp:[LN:HIS1_LACLA] [AC:Q02129] [GN:HISG] [OR:LACTOCOCCUS LACTIS]
[SR:,SUBSPLACTIS:STREPTOCOCCUS LACTIS] [EC:2.4.2.17] [DE:ATP
PHOSPHORIBOSYLTRANSFERASE,] [SP:Q02129] [DB:swissprot] >pir:[LN:D45734] [AC:D45734]
[PN:HisG] [OR:Lactococcus lactis subsp. lactis] [DB:pir2] >gp:[GI:g2565141]
[LN:LLU92974] [AC:U92974:M90760:M90761] [PN:HisG] [GN:hisG] [OR:Lactococcus lactis]
[DB:genpept-bct1] [DE:Lactococcus lactis unknown gene, partial cds, and HisC
(hisC),unknown, HisG (hisG), unknown, HisB (hisB), unknown, HisH (hish),HisA (hisA),
HisF (hisF), HisIE (hisIE), unknown, unknown, LeuA(leuA), LeuB (leuB), LeuC (leuC),
LeuD (leuD), unknown, IlvD(ilvD), IlvB (ilvB), IlvN, IlvC (ilvC), IlvA (ilvA), AldB
(aldB) and aldR (aldR) genes, complete cds.] [NT:phosphoribosyl-ATP synthetase]
[LE:3125] [RE:3751] [DI:direct]

 ORF Name
 NTID
 AAID
 NT AA LengthLength
 Score
 probability

 A17503000985_1207938_c3_2244
 821
 4593
 828
 275
 953
 7.7e-96

Description

gp:[GI:g4433636] [LN:AF029224] [AC:AF029224:AF029225] [PN:NirC] [GN:nirC] [FN:putative nitrite transporter] [OR:Staphylococcus carnosus] [DB:genpept-bct2] [DE:Staphylococcus carnosus nir and nar operons, complete sequences.] [LE:226] [RE:1056] [DI:direct]

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000985_1230437_c1_1528	822	4594	408	135		2.9e-16
Description					L	
sp:[LN:YWBH_BACSU] [AC:P39591] [GN:YDE:HYPOTHETICAL 14.3 KD PROTEIN IN [DB:swissprot] >pir:[LN:S39678] [AC:P39591] [GN:YWBH] [CL:consubtilis] [DB:pir2] >gp:[GI:g413947] subtilis] [DB:genpept-bct1] [DE:B.st[LE:23968] [RE:24354] [DI:complement [AC:Z99123:AL009126] [GN:YWBH] [FN:t0] [DE:Bacillus subtilis complete genom [NT:alternate gene name: ipa-23r] [State	EPR-GALE :S39678:E served hy [LN:BSC abtilis of [LN:BSC abtilis of abtilis of	K INTE F70051 /pothe GENR] genomi GI:e11 [OR:B ion 20	RGENIC PN tical p [AC:X73 c regio 86331:c acillus of 21)	REGION ywbH p protein 3124] [pn (325 g263636 g subti	T] [SP protein HI129 GN:ipa to 33 7] [LN lis]	:P39591] n:hypothetical P7] [OR:Bacillus n-23r] [OR:Bacillus n-23r] [SP:P39591] N:BSUB0020] [DB:genpept-bct1]
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
AI7503000985_12600305_f1_260	823	4595	285	94		
Description	<u> </u>	<u> </u>	J			
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000985_12690706_c3_2230	824	4596	231	76		
Description						
NO-HIT	Marketta against their - To San against their raise and an annual		Comment No. 1001 per la comment		WI Winnessen Hills - J/ I hade	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
AI7503000985_12697136_f2_862	825	4597	873	290	1519	8.1e-156
Description						
pir:[LN:S77609] [AC:S77609] [PN:proprecursor:icaB protein] [GN:icaB] [>gp:[GI:g1161381] [LN:SEU43366] [AC:epidermidis] [DB:genpept-bct2] [DE:Sintercellular adhesion:IcaR, IcaA, I[RE:3134] [DI:direct]	OR:Staph U43366] Staphyloc	nyloco [PN:I coccus	ccus er caB] [0 epider	oidermi SN:icaB rmidis	dis]] [OR: operor	DB:pir2] Staphylococcus nediating
ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	score	probability
A17503000985_12698410_f3_1111	826	4598	150	49		
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000985_127002_£1_259	827	4599	162	53		
Description						

ORF Name	NTID	AAID	<u>NT</u> <u>Length</u>	<u>AA</u> Length	score	probability
A17503000985_1281627_c3_2201	828	4600	1293			1.7e-112
Description	L		JL		الـــــال	
pir:[LN:B69876] [AC:B69876] [PN:ace [OR:Bacillus subtilis] [DB:pir2] >gp [AC:Z99112:AL009126] [GN:ylmB] [FN:um [DE:Bacillus subtilis complete genome [NT:similar to acetylornithine deace states.]	:[GI:e11 nknown] e (secti	185127 [OR:Baion 9	:g26339 acillus of 21):	08] [L subti from	N:BSUB lis] [I 159842]	0009] DB:genpept-bct1] 1to 1807200.]
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000985_129678_£1_53	829	4601	1416	471	1269 2	2.5e-129
Description sp:[LN:YDGF_BACSU] [AC:P96704] [GN:YI						
TRANSPORT PROTEIN IN EXPZ-DINB INTERCOMPIC: [LN:H69782] [AC:H69782] [PN:am: [GN:ydgF] [CL:arginine permease] [OI:gp:[GI:d1020148:g1881368] [LN:AB0014] [SR:Bacillus subtilis (strain:168) DI sequence, 148 kb sequence of the reg: ACID TRANSPORT PERMIASE.] [LE:139917] [CR:Bacillus subtilis] [DB:genpept-bc] [OR:Bacillus subtilis] [DB:genpept-bc] of 21): from 402751 to611850.] [NT [SP:P96704] [LE:203663] [RE:205039] [LN:BSUB004] [AC:Z99107:AL009126] [CDB:genpept-bct1] [DE:Bacillus subtilio813890.] [NT:similar to amino acid [RE:7089] [DI:complement]	ino acid R:Bacill 488] [AC NA] [DB: ionbetwe] [RE:14 003] [AC ct1] [DE :similar [DI:comp GN:ydgF] lis comp	A ABC (Lus sultante) C:AB0000 genperen 35 1293] C:Z9910 C:Bacilot on an	transpo btilis] 1488] [pt-bct1 and 47 [DI:co 06:AL00 llus su mino ac t] >gp: unknown genome	rter ([DB:p GN:ydg] [DE: degre mpleme 9126] btilis id ABC [GI:e1] [OR: (secti	permeas ir2] F] [OR: Bacillu e.] [NT nt] [GN:ydo comple transp 182541: Bacillu on 4 of	se) homolog ydgF] :Bacillus subtilis] us subtilis genome T:PROBABLE AMINO gF] [FN:unknown] ete genome (section porter (permease)] :g2632875] us subtilis] f 21): from 600701
ORF Name	NTID	AAID	<u>NT</u> Lengthl	<u>AA</u> Length	score	probability
AI7503000985_1298202_f2_822	830	4602	1986			2.3e-183
Description pir: [LN:G69848] [AC:G69848] [PN:fruo yjdD] [GN:yjdD] [OR:Bacillus subtili [LN:BSUB0007] [AC:Z99110:AL009126] [OB:genpept-bct1] [DE:Bacillus subtil 1194391to 1411140.] [NT:similar to fi [RE:79575] [DI:direct]	is] [DB: GN:yjdD] lis comp	pir2] [FN:volete o	>gp:[G inknown genome	I:e118] [OR:: (secti	3221:g2 Bacillu on 7 of	2633555] us subtilis] = 21): from
ORF Name	NTID	AAID	<u>NT</u> LengthI	<u>AA</u> Length	score	probability
A17503000985_13089052_c3_2246	831	4603	126	41		
Description	<u> </u>		J			
NO-HIT						
ORF Name A17503000985 1350051 c3 2086		<u>AAID</u> 4604	NT Length!	AA Length	score	probability
Description			الـــــــال		الست	
gp:[GI:g4574118] [LN:AF009415] [AC:AF [OR:Staphylococcus xylosus] [DB:genpetransporter (cudT), putativeregulator dehydrogenase(cudA), and choline dehy [LE:811] [RE:2433] [DI:direct]	ept-bct2 ry prote] [DE: in (cu	Staphy dC), g	lococci lycine	us xylo betain	osus choline ne aldehyde

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000985_1351687_c1_1515	833	4605	906	301	739	3.6e-73
Description gp:[GI:g1644433] [LN:SAU31175] [AC:U dehydrogenase] [GN:ddh] [OR:Staphylo aureus D-specific D-2-hydroxyacid de protein; similar to NAD+-linked D-LD	coccus hydroge	aureus nase (d] [DB:q dh) gei	genpept ne, com	-bct1]	DE:Staphylococcus cds.] [NT:36.7 kDa
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000985_1359635_£3_1248	834	4606	471	156	297	2.5e-26
pir:[LN:F69870] [AC:F69870] [PN:gen [CL:hypothetical protein yklA] [OR:B >gp:[GI:e1181516:g2632036] [LN:BSAJ2 [OR:Bacillus subtilis] [DB:genpept-b between xlyA and ykoR.] [NT:homologo [RE:34555] [DI:direct] >gp:[GI:e1183 [GN:ykzA] [FN:unknown] [OR:Bacillus complete genome (section 7 of 21): f yzzE; similar to general] [LE:187094	acillus 571] [A ct1] [D us to O 336:g26 subtili rom 119	subti C:AJ00 E:Baci smC fr 33670] s] [DB 4391to	lis] [I 2571] llus su om Esch [LN:BS :genper 141114	DB:pir2 [PN:Ykr ubtilis nerichi SUB0007 pt-bct1	A] [Gi A] [Gi 168 ! a coli [] [AC:	N:yknA] [FN:unknown] 56 kb DNA fragment i] [LE:34145] :Z99110:AL009126] :Bacillus subtilis
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000985_1366012_£1_356	835	4607	213	70	81	0.014
Description sp:[LN:PF2R_HUMAN] [AC:P43088] [GN:P(PGF2 ALPHA RECEPTOR)] [SP:P43088] [PN:prostanoid FP receptor] [GN:PTGF sapiens] [SR:, man] [DB:pir2] [MP:1p	DB:swis R:FP]	sprot] [CL:pr	>pir:	[LN:A49	973]	[AC:A49973]
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000985_13678300_f1_40	836	4608	165	54		
Description NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000985_13711588_£2_551	837	4609	285	94	81	0.0019
Description pir: [LN:S53365] [AC:S53365] [PN:muc [SR:, man] [DB:pir2] [MP:11p15.5-11p		(clone	CEL2)]	[GN:M	UC5AC] [OR:Homo sapiens]
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000985_1376926_c3_2008	838	4610	849	282		2.9e-41
Description		<u> </u>	لــــاد			
<pre>gp:[GI:g3127079] [LN:AF061070] [AC:A stutzeri] [DB:genpept-bct2] [DE:Pseu</pre>				_		

gp:[GI:g3127079] [LN:AF061070] [AC:AF061070] [PN:PtxC] [GN:ptxC] [OR:Pseudomonas
stutzeri] [DB:genpept-bct2] [DE:Pseudomonas stutzeri Orf117 (orf117), Orf86 (orf86)
genes, completecds; and ptxABCDE operon, partial sequence.] [NT:putative inner membrane
component of] [LE:3217] [RE:4044] [DI:direct]

ORF Name	NTID	AAID	NT AA Score probability
A17503000985_13796876_£2_710	839	4611	TI80 59
Description NO-HIT			
ORF Name	NTID	AAID	NT AA LengthLength score probability
AI7503000985_13835462_c3_2043	840	4612	1449 482 1386 1.0e-141
Description gp:[GI:e1299584:g3687418] [LN:BLY175 [OR:Bacillus licheniformis] [DB:genp arcC and arcD genes.] [LE:2579] [RE:	ept-bct:	1] [DE	:Bacillus licheniformis arcA, arcB,
ORF Name	NTID	AAID	NT AA score probability
A17503000985_13843910_c1_1637	841	4613	216 71 258 3.9e-22
[OR:Staphylococcus carnosus] [DB:gen [LE:538] [RE:1704] [DI:direct] >gp:[transporter] [GN:narT] [OR:Staphyloc	pept-pat GI:g2529 occus ca Staphylo	t] [DE 9402] arnosu ococcu	[LN:SCU40014] [AC:U40014] [PN:nitrate s] [SR:Staphylococcus carnosus s carnosus nitrate transporter (narT)
ORF Name A17503000985_13866433_f2_661 Description	NTID 842	<u>AAID</u> 4614	NT AA LengthLength score probability 138 45
NO-HIT			
ORF Name	NTID	AAID	NT AA score probability
AI7503000985_13869827_f3_1087	843	4615	2385 794 2176 1.9e-225
[CL:Bacillus probable copper-transpo homology:ATPase transduction domain	rting An homology [DB:pir] nknown] e (secti	TPase y:heav 1] >gp [OR:B ion 18	:[GI:e1186038:g2635863] [LN:BSUB0018] acillus subtilis] [DB:genpept-bct1] of 21): from 3399551to 3609060.]
ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000985_13955288_£1_5	844	4616	165 54
Description NO-HIT			
ORF Name	NTID	AAID	NT AA score probability
A17503000985_14113806_£2_559	845	4617	
Description			J
NO-HIT			

ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	score	probability
A17503000985_14222942_t2_747	846	4618		45		
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000985_1440890_f1_362	847	4619	339	112		
Description						
NO-HIT			 			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000985_14454393_c3_2148	848	4620	144	47		
Description						
NO-HIT					··	
ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	score	probability
A17503000985_14460882_c1_1400	849	4621	177	58	198	7.8e-16
Description		_	_	_		
gp:[GI:g1022726] [LN:SHU35635] [AC:U haemolyticus] [SR:Staphylococcus hae						
[DE:Staphylococcus haemolyticus IS12	_					_
[LE:1101] [RE:1922] [DI:complement]			•	•	-	-
ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	score	probability
ORF Name AI7503000985_14460882_c3_1975	NTID 850	AAID 4622	Length			probability
			Length	Length		
A17503000985_14460882_c3_1975 Description gp:[GI:g1022726] [LN:SHU35635] [AC:U	850 35635]	4622 [PN:un	Length] 177 known]	Length 58 [OR:Sta	242 aphylo	1.7e-20
AI7503000985_14460882_c3_1975 Description gp:[GI:g1022726] [LN:SHU35635] [AC:Uhaemolyticus] [SR:Staphylococcus haemolyticus]	850 335635]	4622 [PN:un	Length	Length 58 [OR:Sta	242 aphylog	1.7e-20 coccus pt-bct1]
A17503000985_14460882_c3_1975 Description gp:[GI:g1022726] [LN:SHU35635] [AC:U	850 335635]	4622 [PN:un	Length	Length 58 [OR:Sta	242 aphylog	1.7e-20 coccus pt-bct1]
Description gp:[GI:g1022726] [LN:SHU35635] [AC:Uhaemolyticus] [SR:Staphylococcus haemolyticus IS12	850 335635]	4622 [PN:un	Length] [177 known] ain=Y17 RF2 gen	[OR:St.	242 aphylog	1.7e-20 coccus pt-bct1]
Description gp:[GI:g1022726] [LN:SHU35635] [AC:Uhaemolyticus] [SR:Staphylococcus haemolyticus IS12 [LE:1101] [RE:1922] [DI:complement]	850 [35635] [molytical [72 ORF1]	4622 [PN:un us str and O	Length [177] known ain=Y17 RF2 gen NT Length	[OR:St.	aphyloc:genpermpletec	coccus pt-bct1] cds.] [NT:ORF1]
Description gp:[GI:g1022726] [LN:SHU35635] [AC:Uhaemolyticus] [SR:Staphylococcus haemolyticus IS12 [LE:1101] [RE:1922] [DI:complement] ORF Name	850 [35635] [molytical [72 ORF1] [NTID]	PN:unus strand O	Length Length 177 known ain=Y17 RF2 gen NT Length	[OR:Standard DB Control Contro	aphyloc:genpermpletec	coccus pt-bct1] cds.] [NT:ORF1]
AI7503000985_14460882_c3_1975 Description gp:[GI:g1022726] [LN:SHU35635] [AC:Union line in the content of the	850 [35635] [molytical [72 ORF1] NTID [851] [35635]	PN:un us str and O AAID 4623 [PN:un	Length [177] known ain=Y17 RF2 gen NT Length [210] known]	[OR:Standard Control C	aphyloc:genpermpleted	coccus pt-bct1] cds.] [NT:ORF1] probability 7.9e-23
A17503000985_14460882_c3_1975 Description gp:[GI:g1022726] [LN:SHU35635] [AC:U haemolyticus] [SR:Staphylococcus hae [DE:Staphylococcus haemolyticus IS12 [LE:1101] [RE:1922] [DI:complement] ORF Name A17503000985_14460882_c3_2098 Description gp:[GI:g1022726] [LN:SHU35635] [AC:U haemolyticus] [SR:Staphylococcus haemolyticus] SR:Staphylococcus haemolyticus SR:Staphylococcus haemolyticus SR:Staphylococcus haemolyticus SR:Staphylococcus haemolyticus SR:Staphylococcus SR:Staphylococ		PN:un and O AAID 4623 [PN:un as structure]	Length [177] known ain=Y17 RF2 gen NT Length [210] known ain=Y17	[OR:Standard Control C	aphylociscore 264 aphylociscore 264 aphylociscore	coccus pt-bct1] cds.] [NT:ORF1] probability 7.9e-23 coccus pt-bct1]
Description gp:[GI:g1022726] [LN:SHU35635] [AC:Uhaemolyticus] [SR:Staphylococcus hae [DE:Staphylococcus haemolyticus IS12 [LE:1101] [RE:1922] [DI:complement] ORF Name A17503000985_14460882_c3_2098 Description gp:[GI:g1022726] [LN:SHU35635] [AC:Uhaemolyticus] [SR:Staphylococcus haemolyticus IS12 [DE:Staphylococcus haemolyticus] [SR:Staphylococcus haemolyticus IS12		PN:un and O AAID 4623 [PN:un as structure]	Length [177] known ain=Y17 RF2 gen NT Length [210] known ain=Y17	[OR:Standard Control C	aphylociscore 264 aphylociscore 264 aphylociscore	coccus pt-bct1] cds.] [NT:ORF1] probability 7.9e-23 coccus pt-bct1]
A17503000985_14460882_c3_1975 Description gp:[GI:g1022726] [LN:SHU35635] [AC:U haemolyticus] [SR:Staphylococcus hae [DE:Staphylococcus haemolyticus IS12 [LE:1101] [RE:1922] [DI:complement] ORF Name A17503000985_14460882_c3_2098 Description gp:[GI:g1022726] [LN:SHU35635] [AC:U haemolyticus] [SR:Staphylococcus haemolyticus] SR:Staphylococcus haemolyticus SR:Staphylococcus haemolyticus SR:Staphylococcus haemolyticus SR:Staphylococcus haemolyticus SR:Staphylococcus SR:Staphylococ		PN:un and O AAID 4623 [PN:un as structure]	Length [177] known] ain=Y17 RF2 gen NT Length [210] known] ain=Y176 RF2 gene	[OR:Standard DB CO CO CO CO CO CO CO C	aphylociscore 264 aphylociscore 264 aphylociscore	coccus pt-bct1] cds.] [NT:ORF1] probability 7.9e-23 coccus pt-bct1]
Description gp:[GI:g1022726] [LN:SHU35635] [AC:Uhaemolyticus] [SR:Staphylococcus hae [DE:Staphylococcus haemolyticus IS12 [LE:1101] [RE:1922] [DI:complement] ORF Name A17503000985_14460882_c3_2098 Description gp:[GI:g1022726] [LN:SHU35635] [AC:Uhaemolyticus] [SR:Staphylococcus haemolyticus IS12 [DE:Staphylococcus haemolyticus] [SR:Staphylococcus haemolyticus IS12		PN:un and O AAID 4623 [PN:un as structure]	Length [177] known ain=Y17 RF2 gen NT Length [210] known ain=Y17	[OR:Standard Control C	aphylociscore 264 aphylociscore 264 aphylociscore	coccus pt-bct1] cds.] [NT:ORF1] probability 7.9e-23 coccus pt-bct1]
Description gp:[GI:g1022726] [LN:SHU35635] [AC:Unamolyticus] [SR:Staphylococcus hae [DE:Staphylococcus haemolyticus IS12 [LE:1101] [RE:1922] [DI:complement] ORF Name A17503000985_14460882_c3_2098 Description gp:[GI:g1022726] [LN:SHU35635] [AC:Unamolyticus] [SR:Staphylococcus haemolyticus] [SR:Staphylococcus haemolyticus] [SR:Staphylococcus haemolyticus IS12 [LE:1101] [RE:1922] [DI:complement]	850 [35635] [molytic: [72 ORF1] NTID [851] [35635] [molytic: [72 ORF1]	PN:un and O AAID 4623 [PN:un as strain and O	Length [177] known ain=Y17 RF2 gen NT Length [210] known ain=Y17 RF2 gen NT Length Length	[OR:Standard Control C	aphylocises score 264 aphylocises aphylo	coccus pt-bct1] cds.] [NT:ORF1] probability 7.9e-23 coccus pt-bct1] cds.] [NT:ORF1]
Description gp:[GI:g1022726] [LN:SHU35635] [AC:Uhaemolyticus] [SR:Staphylococcus hae [DE:Staphylococcus haemolyticus IS12 [LE:1101] [RE:1922] [DI:complement] ORF Name AI7503000985_14460882_c3_2098 Description gp:[GI:g1022726] [LN:SHU35635] [AC:Uhaemolyticus] [SR:Staphylococcus haemolyticus IS12 [LE:1101] [RE:1922] [DI:complement] ORF Name AI7503000985_14460882_c3_2112 ORF Name AI7503000985_14460882_c3_2112 Description	850 [35635] [molytic: [72 ORF1] [851] [35635] [molytic: [72 ORF1] [NTID] [852]	PN:un and O AAID PN:un as tr and O AAID AAID AAID AAID AAID	Length [177] known ain=Y17 RF2 gen NT Length [210] known ain=Y17 RF2 gen NT Length [177]	[OR:Standard Control C	aphyloc:genpermpleted aphyloc:genpermpleted score aphyloc:genpermpleted score	coccus pt-bct1] cds.] [NT:ORF1] probability 7.9e-23 coccus pt-bct1] cds.] [NT:ORF1] probability
Description GRF Name Description GRF Name Description GRF Name DE:Staphylococcus haemolyticus SR:Staphylococcus haemolyticus SR:Staphylococcus haemolyticus SR:Staphylococcus haemolyticus SR:Staphylococcus haemolyticus SR:Staphylococcus haemolyticus GRF Name Description GRF Name Description GRE:Staphylococcus haemolyticus SR:Staphylococcus haemolyticus SR:Staphylococcus haemolyticus SR:Staphylococcus haemolyticus GRE:1922 DI:complement DRF Name	850 [35635] [molytic: 72 ORF1 [851] [35635] [molytic: 72 ORF1 NTID NTID 852 [35635]	PN:un and O AAID PN:un and O AAID AAID AAID AAID AAID AAID	Length [177] known ain=Y17 RF2 gen NT Length [210] known ain=Y17 RF2 gen NT Length [177] Length [177] known]	[OR:Standard Content C	aphylocises aphylo	coccus pt-bct1] cds.] [NT:ORF1] probability 7.9e-23 coccus pt-bct1] cds.] [NT:ORF1] probability 3.4e-19
Description gp:[GI:g1022726] [LN:SHU35635] [AC:Uhaemolyticus] [SR:Staphylococcus hae [DE:Staphylococcus haemolyticus IS12 [LE:1101] [RE:1922] [DI:complement] ORF Name AI7503000985_14460882_c3_2098 Description gp:[GI:g1022726] [LN:SHU35635] [AC:Uhaemolyticus] [SR:Staphylococcus haemolyticus IS12 [LE:1101] [RE:1922] [DI:complement] ORF Name AI7503000985_14460882_c3_2112 ORF Name AI7503000985_14460882_c3_2112 Description		PN:un and O AAID PN:un as strand O AAID AAID AAID AAID AAID AAID AAID AAID AAID	Length [177] known ain=Y17 RF2 gen NT Length [210] known ain=Y17 RF2 gen NT Length [177] known ain=Y17 known ain=Y176	[OR:Standard Content C	aphylocises aphylo	coccus pt-bct1] cds.] [NT:ORF1] probability 7.9e-23 coccus pt-bct1] cds.] [NT:ORF1] probability 3.4e-19 coccus pt-bct1]

ORF Name	NTID	AAID	NT AA LengthLength	probability
A17503000985_14460932_±1_34	853	4625	177 58 208	5.8e-17
Description	•			
gp:[GI:g1022726] [LN:SHU35635] [AC:Uhaemolyticus] [SR:Staphylococcus hae [DE:Staphylococcus haemolyticus IS12 [LE:1101] [RE:1922] [DI:complement]	molytic	cus str	ain=Y176] [DB:genpe	ot-bct1]
ORF Name	NTID	AAID	NT AA LengthLength	probability
AI7503000985_14460932_f2_467	854	4626	177 58 196 1	3e-15
Description				
gp:[GI:g1022726] [LN:SHU35635] [AC:Uhaemolyticus] [SR:Staphylococcus hae [DE:Staphylococcus haemolyticus IS12 [LE:1101] [RE:1922] [DI:complement]	molytic	cus str	ain=Y176] [DB:genper	pt-bct1]
ORF Name	NTID	AAID	NT AA LengthLength	probability
AI7503000985_14460932_f3_1045	855	4627	177 58 218 5	5.9e-18
Description				
<pre>gp:[GI:g1022726] [LN:SHU35635] [AC:U haemolyticus] [SR:Staphylococcus hae [DE:Staphylococcus haemolyticus IS12 [LE:1101] [RE:1922] [DI:complement]</pre>	molytic	cus str	ain=Y176] [DB:genper	ot-bct1]
ORF Name	NTID	<u>AAID</u>	NT AA LengthLength	probability
AI7503000985_14480312_c1_1349	856	4628	1308 435 588 3	6.6e-57
Description				
pir:[LN:F69581] [AC:F69581] [PN:ace acetyltra) acoC] [GN:acoC] [CL:dihy lipoyl/biotin-binding homology] [OR:>gp:[GI:e1182798:g2633132] [LN:BSUB0 E2 component] [GN:acoC] [OR:Bacillus complete genome (section 5 of 21): fyfjI] [LE:77735] [RE:78931] [DI:dire [AC:D78509] [PN:YfjI] [OR:Bacillus s [DB:genpept-bct1] [DE:Bacillus subti [RE:8590] [DI:complement]	drolipe Bacille 005] [A subtil rom 802 ct] >gg ubtilis	pamide us subt AC:Z991 Lis] [D 2821 to p:[GI:d s] [SR:	acetyltransferase: ilis] [DB:pir2] 08:AL009126] [PN:ace B:genpept-bct1] [DE: 1011250.] [NT:alterr 1025206:g2780393] [I Bacillus subtilis (s	etoin dehydrogenase Bacillus subtilis nate gene name: LN:D78509] strain:AC327) DNA]
ORF Name	NTID	AAID	NT AA score	probability
AI7503000985_14490756_£1_295	857	4629	177 58	
Description		-		
NO-HIT		0.00 B	zel an a Arana	
ORF Name	NTID	AAID	NT AA LengthLength	probability
AI7503000985_14492125_f3_1310	858	4630	141 46	
Description				

ORF Name	NTID	AAID LengthLength score probability
AI7503000985_14495712_c1_1593	859	4631 297 98 109 5.0e-06
4.] [PN:pXO1-85] [OR:Bacillus anthracis] rulence plasmid PX01, complete sequence.]
ORF Name	NTID	AAID NT AA score probability
A17503000985_14534387_f3_1231	860	4632 225 74 91 0.00017
haemolytic activity] [OR:Staphylococ	cus lug I A (slu	shA), SLUSH B (slushB), andhemolysin
ORF Name	NTID	AAID NT AA score probability
A17503000985_14552215_c3_2231	861	4633 192 63
Description NO-HIT		
ORF Name	NTID	AAID NT AA score probability
AI7503000985_14562760_f1_359	862	4634 141 46
Description NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000985_14565637_f2_883	863	4635 132 43
<u>Description</u>		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000985_14626432_c2_1687	864	4636 2421 806 1387 7.8e-142
TRANSLOCASE SECA SUBUNIT] [SP:006446 [AC:U97062] [PN:SecA] [GN:secA] [FN:	[DB:s	R:STAPHYLOCOCCUS AUREUS] [DE:PREPROTEIN wissprot] >gp:[GI:g2078390] [LN:SAU97062] on] [OR:Staphylococcus aureus] NCTC 8325 SecA (secA) gene, complete cds.]
ORF Name	NTID	AAID NT AA score probability
A17503000985_14644037_c3_2056	865	4637 [1575] 524 [401 2.8e-40
Description		
sp:[LN:YBJT_ECOLI] [AC:P75822] [GN:Y		R:ESCHERICHIA COLI] [DE:HYPOTHETICAL 53.7 KD

ORF Name	NTID	AAID LengthLength score probability
AI7503000985_14650312_c3_2045	866	4638 1170 389 319 1.2e-28
Description	,	
		AJ007788] [GN:capA] [OR:Bacillus cereus]
[DB:genpept-bct1] [DE:Bacillus cereu		ilvA, capA genes, orf4, orf5 and [LE:2729] [RE:3832] [DI:complement]
orre, partial.) [Nr. related sequence	M24150]	[LE:2729] [RE:3632] [DI:Complement]
ORF Name	NTID	AAID NT AA score probability
A17503000985_14662577_c3_1984	867	4639 537 178
Description		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000985_14720378_c3_2022	868	4640 726 241 292 8.5e-26
Description	<u> </u>	
		al protein aq_928] [GN:aq_928] [OR:Aquifex
		E000714] [AC:AE000714:AE000657] [PN:putative [DB:genpept-bct2] [DE:Aquifex aeolicus
section 46 of 109 of the complete ge		
ORF Name	NTID	AAID LengthLength score probability
A17503000985_14725887_f3 1312	869	4641 183 60
Description		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000985_14729702_c2_1699	870	4642 2031 676 288 7.6e-22
Description		
		[PN:dentin phosphoryn] [OR:Homo sapiens]
[LE:<1] [RE:2367] [DI:direct]	o sapie	ns dentin phosphoryn mRNA, complete cds.]
ORF Name	NTID	AAID NT AA score probability
ORF Name AI7503000985_14742887_c2_1735	NTID 871	AAII) — score probability
		LengthLength probability
AI7503000985_14742887_c2_1735		LengthLength probability
AI7503000985_14742887_c2_1735 Description NO-HIT	871	LengthLength score probability 4643 606 201
A17503000985_14742887_c2_1735 Description		LengthLength probability
AI7503000985_14742887_c2_1735 Description NO-HIT	871	AAID LengthLength score probability AAID NT AA Score probability
AI7503000985_14742887_c2_1735 Description NO-HIT ORF Name AI7503000985_14742937_c1_1449 Description	NTID 872	AAID LengthLength AAID NT AA LengthLength score probability AAID LengthLength score probability 4644 1203 400 94 0.0033
AI7503000985_14742887_c2_1735 Description NO-HIT ORF Name AI7503000985_14742937_c1_1449 Description gp:[GI:g2935567] [LN:AF049856] [AC:A	NTID 872 AF049856]	AAID LengthLength AAID NT AA LengthLength score probability AAID LengthLength score probability 4644 [1203] [400] [94] [0.0033
A17503000985_14742887_c2_1735 Description NO-HIT ORF Name A17503000985_14742937_c1_1449 Description gp:[GI:g2935567] [LN:AF049856] [AC:Apyogenes] [DB:genpept-bct2] [DE:Streens	NTID 872 1F049856]	AAID LengthLength AAID NT AA LengthLength score probability AAID LengthLength score probability 4644 1203 400 94 0.0033 [PN:M protein] [GN:emm] [OR:Streptococcus propagation of the protein of the protein (emm)
AI7503000985_14742887_c2_1735 Description NO-HIT ORF Name AI7503000985_14742937_c1_1449 Description gp:[GI:g2935567] [LN:AF049856] [AC:A	NTID 872 1F049856]	AAID LengthLength AAID NT AA LengthLength score probability AAID LengthLength score probability 4644 1203 400 94 0.0033 [PN:M protein] [GN:emm] [OR:Streptococcus propagation of the protein of the protein (emm)
A17503000985_14742887_c2_1735 Description NO-HIT ORF Name A17503000985_14742937_c1_1449 Description gp:[GI:g2935567] [LN:AF049856] [AC:Apyogenes] [DB:genpept-bct2] [DE:Streens	NTID 872 1F049856]	AAID LengthLength AAID NT AA LengthLength score probability AAID LengthLength score probability 4644 1203 400 94 0.0033 [PN:M protein] [GN:emm] [OR:Streptococcus propagation of the protein of the protein (emm)
Description NO-HIT ORF Name A17503000985_14742937_c1_1449 Description gp:[GI:g2935567] [LN:AF049856] [AC:Apyogenes] [DB:genpept-bct2] [DE:Stregene, partialcds.] [LE:<1] [RE:>403] ORF Name A17503000985_14843762_f3_1171	NTID 872 AF049856]	AAID LengthLength AAID NT AA LengthLength score probability AAID LengthLength score probability 4644 [1203] 400 [94 [0.0033] [PN:M protein] [GN:emm] [OR:Streptococcus propagation of the score probability AAID NT AA score probability
Description NO-HIT ORF Name A17503000985_14742937_c1_1449 Description gp:[GI:g2935567] [LN:AF049856] [AC:Apyogenes] [DB:genpept-bct2] [DE:Stregene, partialcds.] [LE:<1] [RE:>403] ORF Name	NTID 872 AF049856] eptococcu [DI:din	AAID LengthLength AAID NT AA LengthLength score probability AAID LengthLength score probability AAID LengthLength D. 10033 [PN:M protein] [GN:emm] [OR:Streptococcus probability D. 100000000000000000000000000000000000

ORF Name NTID AAID Length ength score probability

A17503000985_14851551_f1_417 | 874 | 4646 | 993 | 330

Description

gp:[GI:g2565150] [LN:LLU92974] [AC:U92974:M90760:M90761] [PN:unknown] [OR:Lactococcus lactis] [DB:genpept-bct1] [DE:Lactococcus lactis unknown gene, partial cds, and HisC (hisC), unknown, HisG (hisG), unknown, HisB (hisB), unknown, HisH (hish), HisA (hisA), HisF (hisF), HisIE (hisIE), unknown, unknown, LeuA(leuA), LeuB (leuB), LeuC (leuC), LeuD (leuD), unknown, IlvD(ilvD), IlvB (ilvB), IlvN, IlvC (ilvC), IlvA (ilvA), AldB (aldB) and aldR (aldR) genes, complete cds.] [NT:ORF14] [LE:10850] [RE:11809] [DI:complement]

ORF Name	NTID	AAID	NT LengthL	<u>AA</u> ength	score	probability
A17503000985_14851713_c2_1650	875	4647	966	321	941	1.4e-94

Description

pir: [LN:D69581] [AC:D69581] [PN:acetoin dehydrogenase E1 component (TPP-dependent alpha subuni) acoA] [GN:acoA] [CL:pyruvate dehydrogenase (lipoamide) alpha chain: thiamine pyrophosphate-binding domain homology] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1182796:g2633130] [LN:BSUB0005] [AC:Z99108:AL009126] [PN:acetoin dehydrogenase El component] [GN:acoA] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 5 of 21): from 802821 to1011250.] [NT:alternate gene name: yfjK] [LE:75688] [RE:76689] [DI:direct] >qp:[GI:d1025208:q2780395] [LN:D78509] [AC:D78509] [PN:YfjK] [OR:Bacillus subtilis] [SR:Bacillus subtilis (strain:AC327) DNA] [DB:genpept-bct1] [DE:Bacillus subtilis YfjG-YfjR genes, complete cds.] [LE:9636] [RE:10637] [DI:complement] >gp:[GI:g2957146] [LN:AF006075] [AC:AF006075] [PN:TPP-dependent acetoin dehydrogenase, E1] [GN:acoA] [OR:Bacillus subtilis] [DB:genpept-bct2] [DE:Bacillus subtilis acetoin dehydrogenase enzyme system gene cluster, ribosomal protein L6-like protein gene, partial cds, TPP-dependentacetoin dehydrogenase, El alpha-subunit (acoA), TPP-dependentacetoin dehydrogenase, E1 beta-subunit (acoB), dihydrolipoamideacetyltransferase (acoC) and dihydrolipoamide dehydrogenase (acoL)genes, complete cds, and regulatory protein (acoR) gene, partialcds.] [NT:alpha subunit of the E1 component of the acetoin] [LE:825] [RE:1826] [DI:direct] >gp:[GI:g2957146] [LN:AF006075] [AC:AF006075] [PN:TPP-dependent acetoin dehydrogenase, E1] [GN:acoA] [OR:Bacillus subtilis] [DB:genpept] [DE:Bacillus subtilis acetoin dehydrogenase enzyme system gene cluster, ribosomal protein L6-like protein gene, partial cds, TPP-dependentacetoin dehydrogenase, El alpha-subunit (acoA), TPP-dependentacetoin dehydrogenase, E1 beta-subunit (acoB), dihydrolipoamideacetyltransferase (acoC) and dihydrolipoamide dehydrogenase (acoL)genes, complete cds, and regulatory protein (acoR) gene, partialcds.] [NT:alpha subunit of the E1 component of the acetoin] [LE:825] [RE:1826] [DI:direct]

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000985_14879667_c2_1781	876	4648	867	288	735	9.7e-73

Description

sp:[LN:PANC_BACSU] [AC:P52998] [GN:PANC] [OR:BACILLUS SUBTILIS] [EC:6.3.2.1] [DE:(PANTOATE ACTIVATING ENZYME)] [SP:P52998] [DB:swissprot] >pir:[LN:H69671] [AC:H69671] [PN:pantothenate synthetase panC] [GN:panC] [CL:pantoate--beta-alanine ligase] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:g1146241] [LN:BACYPIA] [AC:L47709] [PN:pantothenate synthetase] [GN:panC] [FN:pantothenic acid biosynthesis] [OR:Bacillus subtilis] [DB:genpept-bct1] [EC:6.3.2.1] [DE:Bacillus subtilis (clone YAC15-6B) ypiABF genes, qcrABC genes, ypjABCDEFGHI genes, birA gene, panBCD genes, dinG gene, ypmB gene, aspB gene, asnS gene, dnaD gene, nth gene and ypoC gene, completecds's.] [NT:40.8% of identity to the Escherichia coli] [LE:14128] [RE:14988] [DI:direct] >gp:[GI:e1183687:g2634660] [LN:BSUB0012] [AC:Z99115:AL009126] [PN:pantothenate synthetase] [GN:panC] [FN:pantothenate biosynthesis] [OR:Bacillus subtilis] [DB:genpept-bct1] [EC:6.3.2.1] [DE:Bacillus subtilis complete genome (section 12 of 21): from 2195541to 2409220.] [SP:P52998] [LE:156679] [RE:157539] [DI:complement]

OPP Name			NT	AA		
ORF Name	NTID	AAID	Length	Length	score	probability
A17503000985_14880051_f1_116	877	4649	1971	656	2450	1.8e-254
Description						
pir:[LN:C69621] [AC:C69621] [PN:fru subtilis fructose-bisphosphatase: ph [EC:3.1.3.11] [DB:pir1] >gp:[GI:d101 [OR:Bacillus subtilis] [SR:Bacillus [DE:Bacillus subtilis 36kb sequence [LE:9575] [RE:11590] [DI:direct] >gp [AC:Z99124:AL009126] [PN:fructose-1, [OR:Bacillus subtilis] [DB:genpept-b genome (section 21 of 21): from 3999 [LE:127957] [RE:129972] [DI:direct]	nosphoes .1939:g1 subtili between o:[GI:e1 6-bisphoct1] [E	terase 064791 s (str gntZ 184745 osphat C:3.1.	core h] [LN:E ain:168 and trn :g26365 ase] [G 3.11] [nomolog BACGNTZ B) DNA] BY gene G66] [L GN:fbp]	[Y] [OR AND ADDRESS AND ADDRES	:Bacillus subtilis] :D78193] [GN:yydE] enpept-bct1] ing 34 ORFs.] 0021] luconeogenesis] subtilis complete
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000985_14881250_f3_1040	878	4650		332		5.9e-144
Description		L				
<pre>gp:[GI:g1644433] [LN:SAU31175] [AC:U dehydrogenase] [GN:ddh] [OR:Staphylo aureus D-specific D-2-hydroxyacid de protein; similar to NAD+-linked D-LD</pre>	coccus a	aureus nase(d] [DB:g dh) gen	enpept e, com	-bct1]	[DE:Staphylococcus cds.] [NT:36.7 kDa
ORF Name	NTID	AAID	<u>NT</u> Length:	<u>AA</u> Length	score	probability
AI7503000985_14881908_c1_1494	879	4651		70	75	0.042
Description					<u> </u>	
gp:[GI:g4406247] [LN:AF105113] [AC:A [GN:cps19AJ] [OR:Streptococcus pneum pneumoniae type 19A putative oligosa partial cds; UDP-N-acetylglucosamine thymidylyltransferase (cps19AL), dTD dTDP-glucose-4,6-dehydratase (cps19A complete cds; and AliA(aliA) gene, p [DI:direct]	oniae] ccharide -2-epime P-4-kete N), ande	[DB:ge: e repe: erase o-6-de: dTDP-L	npept-b atunit (cps19A oxygluc -rhamno	ct2] [transp K), gl cose-3, ese syn	DE:Stroorter ucose-: 5-epimothase	eptococcus (cps19AJ) gene, 1-phosphate erase(cps19AM), (cps19AO) genes,
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000985_14901578_c3_2235	880	4652		295		3.2e-58
Description						·
pir:[LN:B69772] [AC:B69772] [PN:con [CL:conserved hypothetical protein M >gp:[GI:d1020044:g1881264] [LN:AB001 [SR:Bacillus subtilis (strain:168) D sequence, 148 kb sequence of the reg SIMILAR PRODUCT IN B. SUBTILIS] [LE:	J0449] 488] [A0 NA] [DB ionbetwo	[OR:Bac C:AB00] :genpe een 35	cillus 1488] [pt-bct1 and 47	subtil GN:ydb] [DE: degre	is] [DI O] [OR Bacillu e.] [N]	3:pir2] :Bacillus subtilis] us subtilis genome

>gp:[GI:e1182420:g2632754] [LN:BSUB0003] [AC:Z99106:AL009126] [GN:ydb0] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section

3 of 21): from 402751 to611850.] [NT:similar to hypothetical proteins] [LE:103673]

[RE:104545] [DI:direct]

			NT AA
ORF Name	NTID	AAID	LengthLength score probability
A17503000985_14931501_c2_1790	881	4653	1062 353 802 7.7e-80
Description			
	dum] [SF: AE00118: DB:ger	R:, syp 39:AE00 apept-b	ohilis spirochete] [DB:pir2] 00520] [PN:regulatory protein (pfoS/R)] 0ct2] [DE:Treponema pallidum section 5
ORF Name	NTID	AAID	NT AA score probability
AI7503000985_14979713_c1_1590	882	4654	483 160 251 1.9e-21
Description			
sp:[LN:YORZ_LISMO] [AC:P33385] [OR:I [SP:P33385] [DB:swissprot] >pir:[LN:monocytogenes] [DB:pir2] >gp:[GI:g14] monocytogenes] [SR:Listeria monocytogenes] [DE:Listeria monocytogenes lecithina (plcB) gene complete cds, (ldh) gene [DI:direct]	:F43868] 19647] ogenes (ase, lac	[AC:F [LN:LIS] (strain ctate d	P43868] [PN:ORFZ] [OR:Listeria PACTLDH] [AC:M82881] [OR:Listeria PACTLDH] [DB:genpept-bct1] Pehydrogenase (actA)gene complete cds,
ORF Name	NTID	AAID	NT AA score probability
AI7503000985_15052187_f3_1188	883	4655	7 40
Description			
NO-HIT			
ORF Name A17503000985 15052188 f2 697	NTID	<u>AAID</u>	NT AA score probability
Description	004	14030	
NO-HIT			
ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000985_15126592_f3_1294	885	4657	1086 361 1831 7.0e-189
Description			
pir:[LN:S77610] [AC:S77610] [PN:pro protein] [GN:icaC] [OR:Staphylococo [LN:SEU43366] [AC:U43366] [PN:IcaC] [DB:genpept-bct2] [DE:Staphylococcus	cus epid [GN:ica	lermidi: .C] [OR	s] [DB:pir2] >gp:[GI:g1161382] :Staphylococcus epidermidis]

adhesion:IcaR, IcaA, IcaD, IcaB, and IcaC genes, complete cds.] [LE:3121] [RE:4188]

[DI:direct]

ORF Name	NTID	AAID LengthLength score probability
A17503000985_1537_c1_1639	886	4658 153 50 89 0.00073
[AC:C69783] [PN:NADH dehydrogenase [OR:Bacillus subtilis] [DB:pir2] >gp [GN:ydgI] [OR:Bacillus subtilis] [SR [DB:genpept-bct1] [DE:Bacillus subtiregionbetween 35 and 47 degree.] [NT [RE:146039] [DI:complement] >gp:[GI:[AC:Z99107:AL009126] [GN:ydgI] [FN:u] [DE:Bacillus subtilis complete genoment]	YDGI,] homolog ::[GI:d10 ::Bacillo ::SIMILAI e118254! unknown] ne (sect:	[SP:P96707] [DB:swissprot] >pir:[LN:C69783] ydgI] [GN:ydgI] [CL:nitroreductase] D20152:g1881372] [LN:AB001488] [AC:AB001488] Is subtilis (strain:168) DNA] Ome sequence, 148 kb sequence of the R TO NITROREDUCTASE.] [LE:145410] 5:g2632879] [LN:BSUB0004] [OR:Bacillus subtilis] [DB:genpept-bct1]
ORF Name A17503000985_156387_c1_1367 Description NO-HIT	NTID 887	AAID NT AA score probability LengthLength 4659 138 45
ORF Name A17503000985_15671925_c3_2051 Description NO-HIT	NTID 888	AAID NT AA score probability LengthLength 4660 138 45
ORF Name A17503000985_157093_c2_1947 Description NO-HIT	NTID 889	AAID NT AA score probability LengthLength 48
ORF Name A17503000985_15711457_f3_919 Description NO-HIT	NTID 890	AAID NT AA score probability 4662 201 66
ORF Name A17503000985_157513_c3_2168 Description NO-HIT	NTID 891	AAID NT AA score probability LengthLength 4663 156 51
ORF Name AI7503000985_15752187_f2_664 Description NO_HIT	NTID 892	AAID <u>NT AA</u> score probability 4664 162 53

ORF Name	NTID	AAID NT AA score probability
AI7503000985_157807_c1_1613	893	4665 1584 527 1780 1.8e-183
Description		
		:Y13094] [GN:CTORF585] [OR:Staphylococcus
	ıri mecA	gene, strain K11 (792).] [LE:<1] [RE:1757]
[DI:complement]	S. J. L.	
ORF Name	NTID	AAID NT AA probability LengthLength score
A17503000985_15814001_f1_269	894	4666 153 50
Description		
NO-HIT	·	
ORF Name	NTID	AAID NT AA score probability
A17503000985_158411_c3_2104	895	4667
Description		
sp:[LN:LDH_BACSU] [AC:P13714] [GN:LD [DE:L-LACTATE DEHYDROGENASE,] [SP:P1	H:LCTE] .3714] [[OR:BACILLUS SUBTILIS] [EC:1.1.1.27] DB:swissprot]
ORF Name	NTID	AAID NT AA score probability
AI7503000985 15894527 f3 1275	896	4668 267 88
Description		
NO-HIT		
) TITL
ORF Name	$\underline{\mathtt{NTID}}$	AAID LengthLength score probability
A17503000985_15914762_f1_251	897	4669 579 192
Description		
[OR:Staphylococcus xylosus] [DB:genp	ept-bct	<pre>[PN:putative regulatory protein] [GN:cudC] [DE:Staphylococcus xylosus choline</pre>
transporter (cudT), putativeregulato		
[LE:2604] [RE:3164] [DI:complement]	ıyarogen	ase (cudB) genes, complete cds.] [NT:CudC]
(Latitude of the companion of the compan		
ORF Name	NTID	AAID <u>NT AA</u> score probability LengthLength
A17503000985_16048828_c3_2224	898	4670 261 86
Description		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
AI7503000985_164715_c3_2251	899	4671
Description		
gp:[GI:g4433641] [LN:AF029224] [AC:A		
		t2] [DE:Staphylococcus carnosus nir and nar
operons, complete sequences.] [NT:s1 [LE:5079] [RE:6032] [DI:direct]	milar to	o C-terminus of Esherichia coli CysG.]
[
	Manager and a specific specific	The second section of the second seco
ORF Name	NTID	AAID NT AA score probability
ORF Name A17503000985_16600062_f1_117	NTID 900	
		LengthLength Score probability

<u>NTID AAID NT AA</u> <u>LengthLength</u> score probability A17503000985_16689067_c1_1368 901 4673 597 198 470 1.2e-44	
Sp:[LN:HIS7_LACLA] [AC:Q02134] [GN:HISB] [OR:LACTOCOCCUS LACTIS] [SR:,SUBSPLACTIS:STREPTOCOCCUS LACTIS] [EC:4.2.1.19] [DE:IMIDAZOLEGLYCEROL-PHOSPHATE DEHYDRATASE, (IGPD)] [SP:Q02134] [DB:swissprot] >pir:[LN:G45734] [AC:G45734:C36890] [PN:HisB] [CL:imidazoleglycerol-phosphate dehydratase: imidazoleglycerol-phosphate dehydratase homology] [OR:Lactococcus lactis subsp. lactis] [DB:pir2] >gp:[GI:g2565143] [LN:LLU92974] [AC:U92974:M90760:M90761] [PN:HisB] [GN:hisB] [OR:Lactococcus lactis] [DB:genpept-bct1] [DE:Lactococcus lactis unknown gene, partial cds, and HisC (hisC),unknown, HisG (hisG), unknown, HisB (hisB), unknown, HisH (hish),HisA (hisA), HisF (hisF), HisIE (hisIE), unknown, unknown, LeuA(leuA), LeuB (leuB), LeuC (leuC), LeuD (leuD), unknown, IlvD(ilvD), IlvB (ilvB), IlvN, IlvC (ilvC), IlvA (ilvA), AldB (aldB) and aldR (aldR) genes, complete cds.] [NT:dehydratase] [LE:5869] [RE:6471] [DI:direct]	
NTID AAID NT AA LengthLength AAIT SCORE Probability AAIT LengthLength AAIT LengthLengt	
NTID AAID NT AA probability A17503000985_16802312_c3_2228 903 4675 138 45 Description	

ORF Name

Description NO-HIT

A17503000985_16834377_c2_1861

NTID

904

AAID

4676

147

48

 $\frac{\text{NT}}{\text{Length}} \frac{\text{AA}}{\text{score}}$ probability

			27
ORF Name	NTID	AAID	NT AA LengthLength score probability
AI7503000985_16838207_f1_122	905	4677	288 95 175 2.1e-13
PROTEIN IN DCUB-LYSU INTERGENIC REC >pir:[LN:S56356] [AC:S56356:F65222 intergenic region):hypothetical pro [DB:pir2] >gp:[GI:g536972] [LN:ECOU [DB:genpept-bct1] [DE:Escherichia ominutes.] [NT:ORF_090a] [LE:42913]	GION (090 [PN:hy otein 090 UW93] [AC coli K-12 [RE:4318 [PN:orf, :genpept- genome.]	A)] [Spothet a] [GN chrom 5] [DI hypot bct2]	tical 10.5K protein (dcub-lysu N:yjdJ] [OR:Escherichia coli] 03] [OR:Escherichia coli] mosomal region from 92.8 to 00.1 I:direct] >gp:[GI:g1790569] thetical protein] [GN:yjdJ] [FN:orf; [DE:Escherichia coli K-12 MG1655
[OR:Bacillus subtilis] [DB:genpept- 20 of 21): from 3798401to 4010550.] [RE:189155] [DI:complement] >gp:[GI [GN:yxkD] [OR:Bacillus subtilis] [SI [DB:genpept-bct1] [DE:Bacillus subtilis]	yitT] [O 30020] [A -bct1] [D [NT:sim I:d101238 SR:Bacill tilis gen	R:Baci C:Z991 E:Baci ilar t 7:g178 us sub ome se	hetical protein yxkD] [GN:yxkD] illus subtilis] [DB:pir2] 123:AL009126] [GN:yxkD] [FN:unknown] illus subtilis complete genome (section to hypothetical proteins] [LE:188319] 83243] [LN:D83026] [AC:D83026:D45911] btilis (strain:BGSC 1A1) DNA]
[PN:methylated-DNAprotein-cystein [DB:genpept-bct2] [DE:Haemophilus i	[SP:P4 cotein-cy ne protein [C2] >gp:[ne] [GN:H	4687] steine -cyste GI:g15 I0402] e Rd s	LengthLength 546 181 344 2.6e-31 2] [OR:HAEMOPHILUS INFLUENZAE] [DB:swissprot] >pir:[LN:G64065] e S-methyltransferase homolog] eine S-methyltransferase homology] 573373] [LN:U32723] [AC:U32723:L42023]] [OR:Haemophilus influenzae Rd]
ORF Name AI7503000985_181885_c3_2161	NTID 	<u>AAID</u> 4680	Length ————————————————————————————————————

Description NO-HIT

ORF Name	NTID	AAID	<u>NT</u> LengthL	AA ength ^s	core	probability
AI7503000985_194003_£1_134	909	4681	165 5	54	55	0.025
Description sp:[LN:RK19_GUITH] [AC:078409] [GN:R [DE:CHLOROPLAST 50S RIBOSOMAL PROTEI [LN:AF041468] [AC:AF041468:X14171:X6 [PN:ribosomal protein L19] [GN:rpl19 theta] [DB:genpept-pln2] [DE:Guillar [RE:573] [DI:complement]	N L19] 2349:X5] [OR:C	[SP:078 1511:X1 hlorop]	3409] [D 14504:X5 last Gui	B:swis: 2158:X! llardia	sprot 52912 a the] >gp:[GI:g3602933] :X56806:M7654 7] ta] [SR:Guillardia
ORF Name	NTID	AAID	<u>NT</u> LengthL	AA ength s	core	probability
A17503000985 19531436 c3_2069	910	4682	1365 4		20	4.7e-05
Description		L	ا لــــال		!	
<pre>gp:[GI:g1813493] [LN:BFU64314] [AC:U firmus] [DB:genpept-bct1] [DE:Bacill partial cds.] [NT:similar to Bacillu [DI:direct]</pre>	us firm	us puta	ative hy	drophol	oic p	rotein gene,
ORF Name	NTID	AAID	<u>NT</u> LengthLe	AA ength	core	probability
AI7503000985_19562805_c3_2135	911	4683	213 7	0 6	9	0.036
glycoprotein] [CL:influenza B virus : [MP:segment 6] >gp:[GI:g325221] [LN [SR:Influenza B/Memphis/3/89, cDNA t. B/Memphis/3/89, neuraminidase and NB [RE:315] [DI:direct]	:FLBNAE o viral] [AC:N RNA]	130635] [DB:genp	[OR:Ind	fluen l] [D	za B virus] E:Influenza
ORF Name	NTID	AAID	<u>NT</u> LengthLe	AA ength	core	probability
AI7503000985_19564702_c3_2073	912	4684	477 1	58 3	97	6.3e-37
Description sp:[LN:GSHZ_NICSY] [AC:P30708] [OR:N [EC:1.11.1.9] [DE:GLUTATHIONE PEROXII >pir:[LN:S20501] [AC:S20501] [PN:glute peroxidase] [OR:Nicotiana sylvestris [LN:NS6P229] [AC:X60219] [OR:Nicotiana [DE:N.sylvestris mRNA for 6P229 polymore peroxidases.] [NT:homologous to animal [RE:663] [DI:direct]	DASE HO utathio:] [SR:, na sylve peptide	MOLOG 6 ne pero wood t estris] homolo	SP229,] exidase cobacco] [SR:woo	SP:P30 homolog [DB:pi od toba animal	0708] g] [C ir2] acco] lglut	[DB:swissprot] L:glutathione >gp:[GI:g19739] [DB:genpept-pln1] athione
ORF Name	NTID	AAID	<u>NT</u> LengthLe	AA ength so	core	probability
AI7503000985_19585877_c1_1478	913	4685			34	3.1e-83
Description			: L			
pir:[LN:H69817] [AC:H69817] [PN:amin hydrolase] [OR:Bacillus subtilis] [Di [AC:Z99109:AL009126] [GN:yhaA] [FN:un [DE:Bacillus subtilis complete genome [NT:similar to aminoacylase] [LE:8012]	B:pir2] nknown] e (sect:	>gp:[G OR:Ba ion 6 c	I:e1183 cillus : f 21): :	009:g26 subtili from 99	3334 [1] [1 99501	3] [LN:BSUB0006] DB:genpept-bct1]

ORF Name	NTID	AAID NT AA score probability
A17503000985_19694050_f3_1299	914	4686 147 48
Description		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000985_197090_c2_1776	915	4687 [183] 60
Description NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000985_19709637_c1_1576	916	4688 147 48
Description		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000985_19720642_c2_1710	917	4689 1296 431 525 1.7e-50
desuccinylase] [GN:dapE] [OR:Listeri	la monoc	:AJ007319] [PN:succinyl-diaminopimelate ytogenes] [DB:genpept-bct1] [DE:Listeria genes.] [LE:5533] [RE:6672] [DI:direct]
ORF Name	NTID	AAID NT AA score probability
AI7503000985_197312_c2_1804	918	4690 873 290 813 5.2e-81
1) [SP:007575] [DB:swissprot] 1-dehydrogenase homolog yhdF] [GN:yhhomology] [OR:Bacillus subtilis] [DE [AC:Z99109:AL009126] [GN:yhdF] [FN:ultiple [DE:Bacillus subtilis complete genom [NT:similar to glucose 1-dehydrogenate] >gp:[GI:e1191878:g2226201] [LN:BSY14] [GN:yhdF] [OR:Bacillus subtilis] [DE	>pir:[ndF] [C 3:pir2] unknown] ne (sect ase] [SP 1082] [A 3:genpep to sspB.	L:short-chain alcohol dehydrogenase pgp:[GI:e1182946:g2633280] [LN:BSUB0006] [OR:Bacillus subtilis] [DB:genpept-bct1] ion 6 of 21): from 999501 to1209940.] :007575] [LE:22211] [RE:23080] [DI:direct] C:Y14082] [PN:hypothetical protein] t-bct1] [DE:Bacillus subtilis chromosomal] [NT:Similarity to glucose and ribitol
ORF Name A17503000985 19781305 f1 20	NTID	AAID NT AA score probability LengthLength
	919	4691 123 40
Description NO-HIT	919	4691 123 40

ORF Name	NTID	AAID LengthLength score probability
A17503000985_19929586_c3_2180	921	4693 198 65
Description NO-HIT		
ORF Name	NTID	AAID NT AA score probability
AI7503000985_20081538_f1_158	922	4694 123 40
Description NO-HIT		
ORF Name	NTID	AAID NT AA core probability
A17503000985_20167186_c2_1958	923	4695 156 51 229 4.0e-19
Description gp:[GI:g4096799] [LN:SCU40158] [AC:U [DB:genpept-bct2] [DE:Staphylococcus (orfx)gene, partial cds.] [NT:orfx; [RE:560] [DI:direct]	carnosı	
ORF Name	NTID	AAID NT AA score probability
AT7503000985_20317_c2_1727	924	4696 2088 695 431 8.4e-38
[OR:Bacillus subtilis] [DB:genpept-b (tag3) polypeptide (AA 1-746)] [SP:P >gp:[GI:e1184478:g2636098] [LN:BSUB0 [PN:CDP-glycerol:polyglycerol phosph [OR:Bacillus subtilis] [DB:genpept-b 19 of 21): from 3597091to 3809700.] [LE:78129] [RE:80369] [DI:complement [AC:Z99122:AL009126] [PN:CDP-glycero acid biosynthesis] [OR:Bacillus subt	485] [DI glycerol glycerol rodC:tag ees] >g ct1] [DI 13485] 019] [AG ate] [GI ct1] [DI [NT:alte]] >gp:[Gl:polygl ilis] [I	B:swissprot] >pir:[LN:S06049] ol glycerophosphotransferase, o-phosphotransferase tagF:rodC g-3:tagF] [OR:Bacillus subtilis] gp:[GI:g40100] [LN:BSRODC] [AC:X15200] b:Bacillus subtilis rodC operon.] [NT:rodC [LE:2178] [RE:4418] [DI:direct] C:Z99122:AL009126] w:tagF] [FN:teichoic acid biosynthesis] b:Bacillus subtilis complete genome (section ernate gene name: rodC] [SP:P13485] GI:e1184478:g2636098] [LN:BSUB0019] lycerol phosphate] [GN:tagF] [FN:teichoic DB:genpept] [DE:Bacillus subtilis complete 809700.] [NT:alternate gene name: rodC] ement]
ORF Name	NTID	AAID NT AA score probability
A17503000985_20323403_f3_1320	925	4697 141 46
NO-HIT		
ORF Name	NTID	AAID NT AA probability LengthLength score probability
A17503000985_20335260_c3_2081	926	4698 1881 626 2151 8.6e-223
Description		
[GN:nrdD] [OR:Lactococcus lactis] [D	B:genper naerobic	[PN:anaerobic ribonucleotide reductase] pt-bct2] [DE:Lactococcus lactis anaerobic c ribonucleotide reductase activator protein 67] [RE:2410] [DI:direct]

ORF Name	NTID	AAID NT AA score probability
AI7503000985_20360885_£3_1003	927	4699 1683 560 906 7.3e-91
Description sp:[LN:YHXB_BACSU] [AC:P18159] [GN:Y [DE:PROBABLE PHOSPHOMANNOMUTASE, (PM [AC:C69835:D45868:S18566] [PN:phosp (glpD 3' region)] [GN:YhXB] [OR:Bac >gp:[GI:e1182920:g2633254] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-b 5 of 21): from 802821 to1011250.] [N [LE:203459] [RE:205156] [DI:direct] [AC:Z99109:AL009126] [GN:YhXB] [FN:u [DE:Bacillus subtilis complete genom [NT:similar to phosphomannomutase] [>gp:[GI:e324943:g2226139] [LN:BSY140 [OR:Bacillus subtilis] [DB:genpept-b	HXB] [OIM)] [SP ohomannor illus su oo5] [AG oct1] [DI oce1]	R:BACILLUS SUBTILIS] [EC:5.4.2.8] :P18159] [DB:swissprot] >pir:[LN:C69835] mutase homolog yhxB:hypothetical protein abtilis] [DB:pir2] C:Z99108:AL009126] [GN:yhxB] [FN:unknown] E:Bacillus subtilis complete genome (section ar to phosphomannomutase] [SP:P18159] :e1182932:g2633266] [LN:BSUB0006] [OR:Bacillus subtilis] [DB:genpept-bct1] ion 6 of 21): from 999501 to1209940.] 59] [LE:6779] [RE:8476] [DI:direct] :Y14079] [PN:hypothetical protein] [GN:yhxB] E:Bacillus subtilis chromosomal DNA, region [NT:see EMBL M34393 and Swiss Prot P18159.;
ORF Name A17503000985_20410307_f2_684 Description NO-HIT	NTID 928	AAID NT AA score probability 4700 126 41
ORF Name AI7503000985_20413202_f1_47 Description sp:[LN:YHAD_ECOLI] [AC:P23524] [GN:Y PROTEIN IN RNPB-SOHA INTERGENIC REGI		AAID NT AA score probability 4701 1152 383 961 1.1e-96 R:ESCHERICHIA COLI] [DE:HYPOTHETICAL 39.1 KD 3)] [SP:P23524] [DB:swissprot]
faecalis] [SR:Enterococcus faecalis	plasmid: I17 gene	es for BacA, BacB, ORF3, ORF4, ORF5, ORF6,
ORF Name A17503000985_20484386_f2_470 Description NO-HIT	NTID 931	AAID NT AA score probability 4703 129 42
stutzeri] [DB:genpept-bct2] [DE:Pseud	domonas ron, par	AAID NT AA LengthLength score probability 4704 1011 336 244 1.0e-20 [PN:PtxB] [GN:ptxB] [OR:Pseudomonas stutzeri Orf117 (orf117), Orf86 (orf86) ctial sequence.] [NT:putative binding of:direct]

ORF Name AI7503000985_20509637_c3_2110 Description gp:[GI:g3676414] [LN:AF051917] [AC:A aureus] [DB:genpept-bct2] [DE:Staphy [NT:Orf423] [LE:755] [RE:2026] [DI:d	lococcu		1335 444 311 8.2e-28	
ORF Name AI7503000985_20509637_c3_2186 Description sp:[LN:GNTK_BACLI] [AC:P46834] [GN:G	NTID 934 NTK] [0]	AAID 4706 R:BACI	1545 514 1655 3.1e-170	
[DE:GLUCONOKINASE, (GLUCONATE KINASE [AC:JC2304] [PN:gluconate kinase,:gr [OR:Bacillus licheniformis] [EC:2.7. [LN:BACGNTBL] [AC:D31631] [PN:gluconate kinase,:gr [SR:Bacillus licheniformis] (SR:Bacillus licheniformis (strain:Bacillus licheniformis (strain:Bacillus licheniformis (strain:Bacillus licheniformis DNA for hypothetical proposition [DI:direct]	ntK pro] [D] ate kina GSC5A2)	tein] B:pir2 ase] [DNA]	[GN:gntK] [CL:xylulokinase] [2] >gp:[GI:d1007073:g563951] [GN:gntK] [OR:Bacillus licheniformis] [DB:genpept-bct1] [DE:Bacillus	
ORF Name	NTID	AAID	NT <u>AA</u> LengthLength	
AI7503000985_20515643_f3_1130 Description	935	4707	141 46	
NO-HIT				
ORF Name	NTID	AAID	NT AA score probability	
AI7503000985_20524067_c1_1622	936	4708	639 212 231 2.5e-19	
Description pir: [LN:C70041] [AC:C70041] [PN:constant [OR:Bacillus subtilis] [DB:pir2] >gp [AC:Z99121:AL009126] [GN:yvgV] [FN:ustilis] [DE:Bacillus subtilis complete genome [NT:similar to hypothetical proteins]	:[GI:el: nknown] e (sect:	186036 [OR:B ion 18	6:g2635861] [LN:BSUB0018] Bacillus subtilis] [DB:genpept-bct1] 8 of 21): from 3399551to 3609060.]	

ORF Name NTID AAID NT AA probability

A17503000985_20589568_c1_1392 937 4709 1557 518 282 1.1e-21

Description

sp:[LN:TAGE BACSU] [AC:P13484] [GN:TAGE:RODD:GTAA] [OR:BACILLUS SUBTILIS] [EC:2.4.1.52] [DE: (EC 2.4.1.52) (TEICHOIC ACID BIOSYNTHESIS PROTEIN E)] [SP:P13484] [DB:swissprot] >pir:[LN:S06048] [AC:S06048:F69720] [PN:poly(glycerol-phosphate) alpha-glucosyltransferase, tagE:probable rodD protein:UDP-glucose--polyglycerol phosphate glucosyltransferase tagE] [GN:tagE:rodD] [OR:Bacillus subtilis] [EC: 2.4.1.52] [DB:pir2] [MP:310 degrees] >gp:[GI:g580920] [LN:BSRODC] [AC:X15200] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis rodC operon.] [NT:rodD (gtaA) polypeptide (AA 1-673)] [SP:P13484] [LE:157] [RE:2178] [DI:direct] >gp:[GI:e1184479:g2636099] [LN:BSUB0019] [AC:Z99122:AL009126] [PN:UDP-glucose:polyglycerol phosphate] [GN:tagE] [FN:teichoic acid biosynthesis] [OR:Bacillus subtilis] [DB:genpept-bct1] [EC:2.4.1.52] [DE:Bacillus subtilis complete genome (section 19 of 21): from 3597091to 3809700.] [NT:alternate gene name: rodD, gtaA, gtaD] [SP:P13484] [LE:80369] [RE:82390] [DI:complement] >gp:[GI:e1184479:g2636099] [LN:BSUB0019] [AC:Z99122:AL009126] [PN:UDP-glucose:polyglycerol phosphate] [GN:tagE] [FN:teichoic acid biosynthesis] [OR:Bacillus subtilis] [DB:genpept] [EC:2.4.1.52] [DE:Bacillus subtilis complete genome (section 19 of 21): from 3597091to 3809700.] [NT:alternate gene name: rodD, gtaA, gtaD] [SP:P13484] [LE:80369] [RE:82390] [DI:complement]

 ORF Name
 NTID
 AAID
 NTID
 AAID
 LengthLength
 score
 probability

 A17503000985_2068937_f3_1191
 938
 4710
 1539
 512
 1041
 3.6e-105

Description

pir: [LN:C69676] [AC:C69676:B39096:S16904:I39952:S18269] [PN:alkaline phosphatase, III precursor:alkaline phosphatase B (phoB)] [GN:phoB:phoAIII] [CL:alkaline phosphatase] [OR:Bacillus subtilis] [EC:3.1.3.1] [DB:pir2] >gp:[GI:e1182553:g2632887] [LN:BSUB0004] [AC:Z99107:AL009126] [PN:alkaline phosphatase III] [GN:phoB] [OR:Bacillus subtilis] [DB:genpept-bct1] [EC:3.1.3.1] [DE:Bacillus subtilis complete genome (section 4 of 21): from 600701 to813890.] [NT:alternate gene name: phoAIII] [SP:P19405] [LE:19113] [RE:20501] [DI:complement] >gp:[GI:d1020477:g1945090] [LN:D88802] [AC:D88802] [GN:phoB] [OR:Bacillus subtilis] [SR:Bacillus subtilis (sub_species:Marburg, strain:168, isolate:JH642] [DB:genpept-bct1] [DE:Bacillus subtilis DNA for phoB-rrnE-groESL region, complete cds.] [NT:B. subtilis alkaline phosphatase IIIA; P19405] [LE:6115] [RE:7503] [DI:complement]

NT AΑ ORF Name NTID AAID score probability LengthLength A17503000985 20822287_c1 1574 939 4711 684 227 563 1.6e-54 Description sp:[LN:GNTR BACSU] [AC:P10585] [GN:GNTR] [OR:BACILLUS SUBTILIS] [DE:GLUCONATE OPERON TRANSCRIPTIONAL REPRESSOR (P28 PROTEIN)] [SP:P10585] [DB:swissprot] >pir:[LN:C26190] [AC:C26190:A23537:E69636:S10723] [PN:transcription repressor of gluconate operon gntR:gnt operon regulatory protein] [GN:gntR] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:d1022429:g563933] [LN:AB005554] [AC:AB005554:D45242:D31629] [PN:gluconate operon repressor] [GN:gntR] [OR:Bacillus subtilis] [SR:Bacillus subtilis (strain:BGSC 1A1) DNA] [DB:genpept-bct1] [DE:Bacillus subtilis genomic DNA, 36 kb region between gnt and ioloperons.] [NT:PROSITE; PS00043; HTH GNTR FAMILY; see SWISS PROT] [LE:4516] [RE:5247] [DI:complement] >gp:[GI:g143014] [LN:BACGNT] [AC:J02584:M24505] [PN:gnt repressor] [OR:Bacillus subtilis] [SR:Bacillus subtilis (strain Marburg 168) DNA] [DB:genpept-bct1] [DE:B.subtilis (gluconate operon) gntR, gntK and gntP genes encodinggnt repressor, gluconate kinase and permease, and gntZ gene.] [LE:236] [RE:967] [DI:direct] >qp:[GI:e1184731:q2636552] [LN:BSUB0021] [AC:Z99124:AL009126] [PN:transcriptional regulator (GntR family)] [GN:gntR] [FN:negative regulation of the qluconate operon] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 21 of 21): from 3999281to 4214814.] [SP:P10585] [LE:113345] [RE:114076] [DI:direct] NTAA LengthLength score ORF Name NTID AAID probability A17503000985_20892325_c1_1389 940 4712 156 51 Description NO-HIT NT AΑ ORF Name NTID AAID score probability LengthLength A17503000985 210885 £2 690 941 4713 222 73 Description NO-HIT NT AΑ ORF Name NTID AAID score probability LengthLength A17503000985_2125000_f1_305 942 4714 126 Description NO-HIT NT AA ORF Name NTID AAID score probability LengthLength AI7503000985 2136712 c3 2211 943 4715 1197 398 469 1.5e-44 Description pir:[LN:H69784] [AC:H69784] [PN:chloramphenicol resistance protein homolog ydhL] [GN:ydhL] [CL:Streptomyces lividans chloramphenicol resistance protein] [OR:Bacillus subtilis] [DB:pir2] >qp:[GI:e1182559:q2632893] [LN:BSUB0004] [AC:Z99107:AL009126] [GN:ydhL] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 4 of 21): from 600701 to813890.] [NT:similar to chloramphenicol resistance protein] [LE:24142] [RE:25419] [DI:complement] >gp:[GI:d1020483:g1945096] [LN:D88802] [AC:D88802] [GN:ydhL] [OR:Bacillus subtilis] [SR:Bacillus subtilis (sub_species:Marburg, strain:168, isolate:JH642] [DB:genpept-bct1] [DE:Bacillus subtilis DNA for phoB-rrnE-groESL region, complete cds.]

[NT:S. lividans chloramphenicol resistance protein;] [LE:11144] [RE:12421]

[DI:complement]

ORF Name	NTID	AAID LengthLength score probability
AI7503000985_214026_c2_1767	944	4716 129 42
Description NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000985_2148428_f2_892	945	4717 123 40
Description NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000985_21493827_c2_1719	946	4718 129 42
Description NO-HIT		
ORF Name	NTID	AAID NT AA score probability
AT7503000985_21515707_f3_1084	947	4719 198 65
Description		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000985_21517012_c2_1722	948	4720 411 136 289 1.8e-25
[DB:genpept-bct1] [DE:Bacillus subticds, and YefB (yefB), YefC (yefC), Ye	lis stra eA (yee	PN:YeeE] [GN:yeeE] [OR:Bacillus subtilis] ain 168 trpC2 YefA (yefA) gene, partial A), YeeB (yeeB), YeeC(yeeC), YeeD (yeeD), cds.] [LE:8308] [RE:9417] [DI:complement]
ORF Name	NTID	AAID NT AA score probability
AI7503000985_21523377_f3_1042	949	4721 171 56
Description NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000985_21563751_f3_1315	950	4722 237 78
Description NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000985_21579561_c2_1962	951	4723 123 40
Description NO-HIT		

ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	score	probability
AT7503000985_21604040_c1_1522	952	4724		675		4.4e-132
Description sp:[LN:FEOB_METJA] [AC:Q57986] [GN:M. IRON TRANSPORT PROTEIN B HOMOLOG] [S: [AC:F64370] [PN:ferrous iron transport B:translation elongation factor Tu homological [MP:REV504509-502503] >gp:[GI:g1591: iron transport protein B (feoB)] [GN:genpept-bct2] [DE:Methanococcus genome.] [NT:similar to SP:P33650 PIII [DI:complement]	P:Q57980 ort proposomology 272] [Li :MJ0566] jannasci	6] [DB tein B] [OR:I N:U6756] [OR:I hii se	:swissp] [CL:f Methano 05] [AC Methano ction 4	rot] > errous coccus :U6750 coccus 7 of 1	pir:[I iron janna 5:L771 janna .50 of	N:F64370] transport protein schii] [DB:pir2] .17] [PN:ferrous schii] the complete
ORF Name AI7503000985_21664126_c2_1712 Description gp:[GI:e1299583:g3687417] [LN:BLY175] [GN:arcB] [OR:Bacillus licheniformis] licheniformis arcA, arcB, arcC and a] [DB:ge	enpept	4] [PN:	360 ornith	1138 line ca 1.3.3]	[DE:Bacillus
ORF Name AI7503000985_21674062_c1_1615 Description NO-HIT	<u>NTID</u> 954	<u>AAID</u> 4726	NT LengthI	AA Length 44	score	probability
ORF Name AI7503000985_21678217_c2_1764 Description NO-HIT	<u>NTID</u> 955	<u>AAID</u> 4727	NT LengthI	<u>AA</u> Length 40	score	probability
ORF Name AI7503000985_21687963_c1_1426 Description gp:[GI:g4980875] [LN:AE001717] [AC:AF [GN:TM0371] [OR:Thermotoga maritima] 29 of 136 of the complete genome.] [P [LE:2550] [RE:3008] [DI:direct] >gp: [PN:arginine repressor] [GN:argR] [FR [OR:Thermotoga maritima] [DB:genpept] [LE:1] [RE:459] [DI:direct]	[DB:ger NT:simi] [GI:e148 N:regula	:AE0005 npept-klar to 39641:gation o	512] [Proct2] [I GB:M278 g5102818 of argin	N:argi DE:The 869 SP B] [LN	nine r rmotog :P1789 :TMA13	a maritima section 3 PID:142450] 2286] [AC:AJ132286] hesis genes]
ORF Name AI7503000985_21759718_f1_181 Description gp:[GI:g4049770] [LN:AF063866] [AC:AF family] [GN:MSV254] [OR:Melanoplus sa [DE:Melanoplus sanguinipes entomopoxy [DI:direct]	anguinip	4729 [PN:Opes ent	ORF MSV2	98 254 le virus]	الــــا ucine [DB:g	enpept-vrl]

ORF Name	NTID	AAID NT AA score probability
A17503000985_21774087_c3_2111	958	4730 732 243 336 1.8e-30
Description	L	
sp:[LN:Y882_HAEIN] [AC:P44068] [GN:HPROTEIN HI0882] [SP:P44068] [DB:swister [PN:hypothetical protein HI0882] [OR:pgp:[GI:g1573906] [LN:U32770] [AC:U3276] [DE:Haemophilus influenzae Rd section HI0882] [DE:Ha	sprot] 1:Haemop 2770:L4 philus on 85 of	>pir:[LN:E64015] [AC:E64015] hilus influenzae] [DB:pir2] 2023] [PN:H. influenzae predicted coding influenzae Rd] [DB:genpept-bct2]
ORF Name	NTID	AAID NT AA score probability
AI7503000985_22275052_f1_248	959	4731 135 44
Description NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000985_22296927_f1_418	960	4732 168 55
Description NO-HIT		
ORF Name	NTID	AAID NT AA score probability
AI7503000985_22304578_c2_1792	961	4733 138 45
Description		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000985_22345265_f1_27	962	4734 189 62
Description NO-HIT		
NO-HII		
ORF Name	NTID	AAID NT AA score probability LengthLength
A17503000985_22380343_f2_827	963	4735 1983 660 363 2.3e-30
[EC:3.5.1.28] [DE:AUTOLYSIN, (N-ACET [DB:swissprot] >pir:[LN:A38109] [AC: [DB:pir2] >gp:[GI:g829194] [LN:STRHY hydrolase] [OR:Enterococcus faecalis	YLMURAMO A38109] DROLA]] [SR:St	PN:autolysin] [OR:Enterococcus faecalis]
ORF Name	איזייט	AAID NT AA score probability
ORF Name	NTID	AAID LengthLength score probability
A17503000985_2242136_c3_2048	964	4736 684 227 583 1.2e-56
Description gp:[GI:e303881:g1850807] [LN:CPCPEAA [OR:Clostridium perfringens] [DB:gen genes.] [LE:2477] [RE:2932] [DI:dire	pept-bct	71844] [PN:putative transposase] [DE:C.perfringens uapC, cpe, and nadC

ORF Name A17503000985_22453186_f3_989 Description	<u>NTID</u>	AAID NT AA score probability 4737 159 52
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000985_22460302_c1_1511	966	4738 756 251 334 3.0e-30
YwpF homolog, single-strand binding p	pept-bct roteinde rotein h (sceE) g	[DE:Staphylococcus carnosus chydrase homolog (fabZ) gene, partial cds, nomolog (ssb), SceD precursor (sceD),SceA genes, completecds, and TenA homolog (tenA)
ORF Name	NTID	AAID NT AA score probability
A17503000985_22656300_£2_881	967	4739 1542 513 1508 1.2e-154
	5) DNA]	[DB:genpept-bct1] [DE:Bacillus halodurans] [DB:genpept-bct1] [DE:Bacillus halodurans [Lete cds.] [NT:unknown] [LE:4328] [RE:5830]
ORF Name	NTID	AAID NT AA score probability
A17503000985_22735877_±3_1047 Description NO-HIT	968	4740 141 46
ORF Name A17503000985_23438751_f2_550 Description NO-HIT	NTID 969	AAID NT AA score probability LengthLength 75
ORF Name A17503000985_234432_c1_1440 Description NO-HIT	NTID 970	AAID NT AA score probability 4742 150 49
ORF Name AI7503000985_23445387_c3_1998 Description NO-HIT	NTID 971	AAID NT AA score probability LengthLength 4743 195 64

ORF Name	NTID AAID NT AA score probability
AI7503000985 23468942_f1_79	972 4744 1101 366 928 3.4e-93
Description	
pir: [LN:A69984] [AC:A69984] [PN:end [CL:thermophilic aminopeptidase I al >gp: [GI:e1184131:g2635347] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-b 15 of 21): from 2795131to 3013540.] [RE:155284] [DI:complement] >gp: [GI: [PN:hypothetical protein] [GN:ysdC]	do-1,4-beta-glucanase homolog ysdC] [GN:ysdC] lpha chain] [OR:Bacillus subtilis] [DB:pir2] 0015] [AC:Z99118:AL009126] [GN:ysdC] [FN:unknown] bct1] [DE:Bacillus subtilis complete genome (section [NT:similar to endo-1,4-beta-glucanase] [LE:154199] 101:65304:g1770012] [LN:BSZ75208] [AC:Z75208] [OR:Bacillus subtilis] [DB:genpept-bct1] 109bp.] [NT:homology to celA of Clostridium 1708] [DI:direct]
ORF Name	$rac{ ext{NTID}}{ ext{AAID}} = rac{ ext{NT}}{ ext{Length Length}} rac{ ext{score}}{ ext{probability}}$
A17503000985_23475251_c2_1952	973 4745 141 46
Description	
NO-HIT	
ORF Name	NTID AAID NT AA score probability
A17503000985_23475325_c1_1479	974 4746 159 52
Description	
NO-HIT	
ORF Name	NTID AAID NT AA score probability
A17503000985_23476503_f3_1228	975 4747 189 62
Description	
NO-HIT	
ORF Name	$rac{ ext{NTID}}{ ext{Length}} rac{ ext{AA}}{ ext{Length}} rac{ ext{Score}}{ ext{probability}}$
AI7503000985_23476702_c1_1496	976 4748 882 293 595 6.6e-58
Description	
[DB:swissprot] >pir:[LN:S39679] [AC:ywbI:protein ipa-24d] [GN:ywbI] [CL[OR:Bacillus subtilis] [DB:pir2] >gp [OR:Bacillus subtilis] [DB:genpept-b [SP:P39592] [LE:24460] [RE:25365] [D [AC:Z99123:AL009126] [GN:ywbI] [FN:umather [CE:Bacillus subtilis complete genome	WBI:IPA-24D] [OR:BACILLUS SUBTILIS] GULATOR IN THIK-EPR INTERGENIC REGION] [SP:P39592] S39679:G70051] [PN:transcription regulator homology E:probable transcription regulator lsyR] D:[GI:g413948] [LN:BSGENR] [AC:X73124] [GN:ipa-24d] Doct1] [DE:B.subtilis genomic region (325 to 333).] DI:direct] >gp:[GI:e1186330:g2636366] [LN:BSUB0020] DINKNOWN] [OR:Bacillus subtilis] [DB:genpept-bct1] The (section 20 of 21): from 3798401to 4010550.] Dimilar to] [SP:P39592] [LE:132594] [RE:133499]
ORF Name	NTID AAID NT AA score probability
A17503000985_23479702_c3_2053	977 4749 273 90
Description	
NO UTT	

ORF Name	NTID AAID NT AA score probability
A17503000985_23480467_c1_1549	978 4750 912 303 862 3.4e-86
[OR:Bacillus subtilis] [DB:pir2] >gp [AC:Z99112:AL009126] [GN:ylpA] [FN:u [DE:Bacillus subtilis complete genom [NT:similar to L-serine dehydratase] >gp:[GI:e323528:g2337815] [LN:BSY139	37] [AC:Y13937] [PN:putative YhaP protein] subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis
ORF Name A17503000985_235205_c2_1845 Description	NTID AAID NT AA score probability 979 4751 138 45
NO-HIT	
ORF Name	NTID AAID NT AA score probability LengthLength
AI7503000985_23553275_f3_1064	980 4752 126 41
<u>Description</u>	
NO-HIT	
ORF Name	NTID AAID NT AA score probability
A17503000985_23556338_c1_1568	981 4753 303 100 170 7.2e-13
[AC:A69764] [PN:conserved hypothetic [DB:pir2] >gp:[GI:e1182354:g2632688] [FN:unknown] [OR:Bacillus subtilis] genome (section 3 of 21): from 40275 [SP:P94425] [LE:36112] [RE:36399] [DI [LN:D50453] [AC:D50453] [GN:ycnE] [OX (strain:168 trpC2) DNA] [DB:genpept-1]	CNE] [OR:BACILLUS SUBTILIS] [DE:HYPOTHETICAL 10.9 GION] [SP:P94425] [DB:swissprot] >pir:[LN:A69764] cal protein ycnE] [GN:ycnE] [OR:Bacillus subtilis] [LN:BSUB0003] [AC:Z99106:AL009126] [GN:ycnE] [DB:genpept-bct1] [DE:Bacillus subtilis complete 1 to611850.] [NT:similar to hypothetical proteins] I:complement] >gp:[GI:d1009654:g1805457] R:Bacillus subtilis [SR:Bacillus subtilis bct1] [DE:Bacillus subtilis DNA for 25-36 degree n, complete cds.] [LE:118515] [RE:118802]
ORF Name	NTID AAID NT AA score probability
A17503000985_23593800_c1_1493	982 4754 942 313 141 8.3e-08
subtilis] [DB:pir2] >gp:[GI:d1020110 [OR:Bacillus subtilis] [SR:Bacillus subtilis] [DE:Bacillus subtilis genome sequence degree.] [NT:FUNCTION UNKNOWN.] [LE:>gp:[GI:e1182488:g2632822] [LN:BSUB06]	O03] [AC:Z99106:AL009126] [GN:ydeJ] [FN:unknown]

			NT AA	
ORF Name	NTID	AAID	Length Length score	probability
A17503000985_23595262_t2_695	983	4755	123 40	
Description				
NO-HIT				
ORF Name	NTID	AAID	NT AA LengthLength score	probability
AI7503000985_23595312_f3_948	984	4756	132 43	
Description				
NO-HIT				
ORF Name	NTID	AAID	NT AA LengthLength	probability
AI7503000985_23600412_f3_960	985	4757	1242 413 546 1	.0e-52
Description				
] [CL:bicyclomycin resistance protei >gp:[GI:d1020154:g1881374] [LN:AB001 [SR:Bacillus subtilis (strain:168) Disequence, 148 kb sequence of the reg BICYCLOMYCIN RESISTANCE PROTEIN.] [L >gp:[GI:e1182547:g2632881] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-b 4 of 21): from 600701 to813890.] [NT [LE:12656] [RE:13864] [DI:direct]	488] [A0 NA] [DB ionbetwe E:146860 004] [A0 ct1] [DI	C:AB00 genpe een 35 D] [RE C:Z991 E:Baci	1488] [GN:ydgK] [OR: pt-bct1] [DE:Bacillu and 47 degree.] [NT ::148068] [DI:direct] 07:AL009126] [GN:ydg llus subtilis comple	Bacillus subtilis] s subtilis genome c:SIMILAR TO [K] [FN:unknown] te genome (section
ORF Name	NTID	AAID	NT AA LengthLength score	probability
AI7503000985_23600752_f1_215	986	4758	240 79 73 0	.014
Description				
pir:[LN:S69592] [AC:S69592] [PN:hypoterevisiae] [DB:pir2] [MP:4R]	othetica	al pro	tein YDR509w] [OR:Sa	ccharomyces
ORF Name	NTID	AAID	NT AA LengthLength	probability
A17503000985_23602015_c2_1892	987	4759		.7e-34
Description				
sp:[LN:YBBL_ECOLI] [AC:P77279] [GN:Y:TRANSPORTER ATP-BINDING PROTEIN YBBL [AC:A64780] [PN:probable ABC-type t:ATP-binding protein ybbL] [GN:ybbL] coli] [DB:pir2] >gp:[GI:g1773172] [LI:DB:genpept-bct1] [DE:Escherichia coli] [NT:hypothetical protein] [LE:95367] [LN:AE000155] [AC:AE000155:U00096] [GN:ybbL] [FN:putative transport; Note that is a simple of the color of th	SP:P7 [SP:P7 [CL:ATF [CL:ATF [CU826] [SE:960] [RE:960]	77279] prot P-bind 664] [ces 9	[DB:swissprot] >pir ein ybbL:probable AB ing cassette homolog AC:U82664] [OR:Esche to 11 genomic sequen DI:direct] >gp:[GI:g	:[LN:A64780] C transporter, y] [OR:Escherichia richia coli] ce.] 1786698]

ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	score	probability
A17503000985_23604052_f1_209	988	4760		105		0.0020
Description sp:[LN:YE1A_METJA] [AC:P81328] [GN:M [DE:HYPOTHETICAL PROTEIN MJ1417.1] [[LN:U67582] [AC:U67582:L77117] [PN:M [GN:MJ1417.1] [OR:Methanococcus jann jannaschii section 124 of 150 of the by GeneMark; putative] [LE:4562] [RE	SP:P8132 . jannas aschii] complet	8] [D] schii p [DB:ge se geno	3:swiss predict enpept- ome.] (sprot] :ed cod :bct2]	>gp:[G ing re [DE:Me	I:g2826408] gion MJ1417.1] thanococcus
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000985_23620252_c1_1606 Description NO-HIT	989	4761		45		
ORF Name A17503000985_23625005_c3_2103 Description NO-HIT	<u>NTID</u> 990	<u>AAID</u> 4762	NT Length	AA Length 45	score	probability
ORF Name AI7503000985_23631311_c2_1820 Description NO-HIT	<u>NTID</u> 991	<u>AAID</u> 4763	<u>NT</u> Length	AA Length 98	score	probability
ORF Name A17503000985_23635926_t2_538 Description NO-HIT	<u>NTID</u> 992	AAID 4764	<u>NT</u> Length	AA Length 47	score	probability
ORF Name	NTID	AAID	NT	AA	score	probability
A17503000985_23636343_c1_1383 Description sp:[LN:PHNE_ECOLI] [AC:P16683:Q47479 [DE:PHOSPHONATES TRANSPORT SYSTEM PEI [SP:P16683:Q47479:P76792:Q47716] [DB [AC:F35718:A42732:S56332:S56331:G652:b4103] [GN:phnE] [CL:phnE protein] [LN:ECOPHNAQ] [AC:J05260] [OR:Escher:[DB:genpept-bct1] [DE:E.coli psiD locAthrough Q, complete cds.] [NT:phnE]	:P76792: RMEASE P:swisspr 19:F6521 [OR:Eschichia co	Q47716 ROTEIN ot] >r 9] [F erichi li] [S aining	[GN: PHNE] pir:[LN PN:phnE a coli GR:E.co	PHNE] ::F3571 prote] [DB:: li (st.: phospho	[OR:ES 8] in:hyp pir1] rain B onate	Othetical protein ogp:[GI:g147198] DNA] uptake (phn) genes
ORF Name A17503000985_23642942_c3_2087 Description NO-HIT	NTID 994	<u>AAID</u> 4766	NT Length	AA Length	score	probability

ORF Name	NTID	AAID	NT AA Length Length score	probability
A17503000985_23649187_c3_2050	995	4767		.2e-15
Description				
<pre>gp:[GI:g4049717] [LN:AF063866] [AC:A [GN:MSV156] [OR:Melanoplus sanguinip sanguinipes entomopoxvirus, complete</pre>	es entom	nopoxv	irus] [DB:genpept-vr	[DE:Melanoplus
ORF Name	NTID	AAID	NT AA LengthLength	probability
A17503000985_23672518_f3_1031	996	4768	1011 336 529 6	.5e-51
Description				
pir:[LN:F69659] [AC:F69659] [PN:mol [OR:Bacillus subtilis] [DB:pir2] >gp [AC:Z99111:AL009126] [PN:molybdopter subtilis] [DB:genpept-bct1] [DE:Baci from 1394791to 1603020.] [LE:100770] [LN:AF012285] [AC:AF012285:AF012284: MoeB] [GN:moeB] [OR:Bacillus subtili mobA-nprE gene region.] [NT:member o [RE:2277] [DI:direct]	:[GI:e11 in biosy llus sub [RE:101 U51911] s] [DB:g	.85017 Inthesi otilis .789] [PN:mo genpept	g2633798] [LN:BSUB0 is protein] [GN:moeE complete genome (se [DI:direct] >gp:[GI: plybdopterin biosynt t-bct2] [DE:Bacillus	[008] [OR:Bacillus ction 8 of 21): [g3282111] [hesis protein subtilis
ORF Name	NTID	AAID	<u>NT AA</u> LengthLength	probability
AI7503000985_23680300_f1_330	997	4769	948 315	
Description NO-HIT				
ORF Name	NTID	AAID	<u>NT AA</u> LengthLength	probability
A17503000985_23710811_c1_1425	998	4770	159 52	
Description NO-HIT				
ORF Name	NTID	AAID	NT AA LengthLength	probability
A17503000985_23828253_f1_81	999	4771	156 51	
Description				
NO-HIT				
ORF Name	NTID	AAID	NT AA LengthLength	probability
A17503000985_23866552_f2_815	1000	4772	141 46	
Description	(
NO-HIT				
ORF Name	NTID	AAID.	NT AA LengthLength	probability
AI7503000985_23868887_c2_1828	1001	4773	<u>189 62 </u>	
Description				

ORF Name AI7503000985_24015687_c1_1476 Description gp:[GI:g4574120] [LN:AF009415] [AC:A dehydrogenase] [GN:cudA] [OR:Staphyl [DE:Staphylococcus xylosus choline t (cudC), glycine betaine aldehyde deh genes, complete cds.] [NT:CudA] [LE:	F009415] ococcus ransporte	[PN:g xylosu er (cu se(cud	plycine us] [DB udT), p lA), an	betaine a genpept-boutativereg	1.1e-22 ldehyde ct2]	protein
ORF Name			NT Length		e proba	ability
A17503000985_24019026_f1_390	1003	4775	156	51		
Description NO-HIT						
ORF Name AI7503000985_24022177_c1_1575 Description sp:[LN:GNTP_BACSU] [AC:P12012] [GN:G [SP:P12012] [DB:swissprot] >pir:[LN:gntP] [GN:gntP] [CL:D-serine permea >gp:[GI:d1022427:g563931] [LN:AB0055 permease] [GN:gntP] [OR:Bacillus subticoloperons.] [NT:homologs are found [DI:complement] >gp:[GI:g143016] [LN [OR:Bacillus subtilis] [SR:Bacillus [DB:genpept-bct1] [DE:B.subtilis (glencodinggnt repressor, gluconate kin [RE:3876] [DI:direct] >gp:[GI:e11847 [PN:gluconate permease] [GN:gntP] [F [DB:genpept-bct1] [DE:Bacillus subtilis] [SR:genpept-bct1] [DE:Bacillus subtilis] [SR:genpept-bct1] [DE:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis] [SP:P12012] [LE:	NTP] [OR A26190] se] [OR: I 54] [AC: I tilis] [S lis genor in E. col: BACGNT] subtilis uconate case and I 33:g26369 N:gluconalis complete complete case and I selected to the case and I selected to the complete case and I selected to the case and I selecte	:BACIL [AC:A2 Bacill AB0055 SR:Bac mic DN li and [AC:J (stra operon permea 554] [ate ut lete g	LUS SU 6190:D us sub 554:D45 illus IA, 36 IH. in 102584: in Mar in Mar LN:BSU ilizat enome	BTILIS] [D. 69636] [P. tilis] [DB 242:D31629 subtilis (kb region fluenzae;] M24505] [P. burg 168) , gntK and d gntZ general B0021] [AC ion] [OR:B (section 2	I.le-15 E:GLUCON N:glucon pir2] [PN:glustrain:Bo	ATE PERMEASE] ate permease uconate GSC 1A1) DNA] gnt and 7] [RE:2953] se] nes 2530] AL009126] subtilis]
ORF Name AI7503000985_24070155 t2 878	<u> </u>	AAID <u>1</u>		<u>AA</u> Length scor	proba	ability
Description pir: [LN:A69849] [AC:A69849] [PN:hyp subtilis] [DB:pir2] >gp:[GI:e1183223 [GN:yjdF] [FN:unknown] [OR:Bacillus complete genome (section 7 of 21): f	othetical g2633557 subtilis	l prot 7] [LN] [DB:	ein yj :BSUB0 genpep	dF] [GN:yjo 007] [AC:Z: t-bct1] [Di	JF] [OR: 99110:AL0 ::Bacill	Bacillus 009126] us subtilis

[DI:direct]

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probabil	ity
AI7503000985_24087760_c2_1742	1006	4778	903	300	665	2.5e-65	
Description pir:[LN:F69997] [AC:F69997] [PN:hyp subtilis] [DB:pir2] >gp:[GI:el184177 [GN:ytnM] [FN:unknown] [OR:Bacillus complete genome (section 15 of 21): [DI:complement] >gp:[GI:el185801:g26 [FN:unknown] [OR:Bacillus subtilis] genome (section 16 of 21): from 2997 >gp:[GI:g2293257] [LN:AF008220] [AC: subtilis] [DB:genpept-bct2] [DE:Baci [NT:similar to a hypothetical protei	:g26353 subtili: from 27 35412] [DB:gen] 771to 33 AF00822	93] [Lis] [DB 95131to [Lin:BS] [Pept-bo 213410] [Pin btilis]	N:BSUBC :genper 0 30135 UB0016] ct1] [LE: :YtnM] rrnB-c	0015] ot-bct1 540.] [AC:2 DE:Baci :99] [F [GN:yt] dnaB ge	[AC: Z95 L] [DE: [LE: 202 Z99119: illus s RE: 1003 cnM] [C	Bacillus s Bacillus s 2739] [RE:2 AL009126] Subtilis co DR:Bacillus region.]	.26] subtilis :03641] [GN:ytnM] omplete olement]
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	score	probabil	ity
AI7503000985_24095327_c2_1770	1007	4779		44]		
Description NO-HIT							
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probabil	ity
A17503000985_24101701_c1_1485	1008	4780		513		2.9e-158	
<pre>gp:[GI:d1039113:g4514332] [LN:AB0133 [SR:Bacillus halodurans (strain:C-12 C-125 yesT and comEC genes, partial [DI:direct]</pre>	5) DNA]	[DB:ge	enpept-	bct1]	[DE:Ba	cillus hal	odurans
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probabil	ity
A17503000985_24220260_c3_2109	1009	4781	1056	351	121	0.00034	
Description gp:[GI:e316518:g2230824] [LN:DDSTATF [FN:regulates stalk cell differentia [DB:genpept-inv1] [DE:D.discoideum m [RE:2630] [DI:direct]	tion] [(OR:Dict	yostel	ium di	scoide	um]	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probabil	ity
AI7503000985_24220290_f2_582	1010	4782	1482	493	935	6.2e-94	
Description sp:[LN:ALDA_ECOLI] [AC:P25553] [GN:A [DE:ALDEHYDE DEHYDROGENASE A, (LACTA							
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probabil	ity
A17503000985_24224037_c2_1825 Description NO-HIT	1011	4783	-	47	•		
ORF Name A17503000985_24225015_f3_1098 Description	NTID 1012	<u>AAID</u> 4784	NT Length	AA Length 55	score	probabil	ity

ODE W		NT AA
ORF Name	NTID	AAID LengthLength score probability
A17503000985_24225015_f3_1127	1013	4785 141 46
Description		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000985_24226577_c3_2184	1014	4786 834 277 153 9.4e-11
Description		
RESISTANCE OPERON REGULATORY PROTEIN [AC:A32227] [PN:hypothetical protein [OR:Bacillus sp.] [DB:pir2] >gp:[GI:GI:PN:regulatory protein] [GN:merR] [OI:Bacillus megaterium ORF2, ORF3, GI:direct] >gp:[GI:g1129093] [LN:AF:[PN:mercury resistance operon negatives]	[SP:P2 n 1 (mer e301549: R:Bacill DRF4, me 138877] ve regul	us megaterium] [DB:genpept-bct1] erR and merA genes.] [LE:574] [RE:972]
ORF Name	NTID	AAID NT AA score probability
AI7503000985_24250177_c1_1630	1015	4787 714 237 653 4.7e-64
Description		
carnosus] [DB:genpept-pat] [DE:Sequer product] [LE:9374] [RE:9949] [DI:dire [AC:AF029224:AF029225] [PN:NarJ] [GN	nce 1 fr ect] >gr :narJ] carnosu	OR:Staphylococcus carnosus] s nir and nar operons, complete sequences.]
ORF Name	NTID	AAID NT AA score probability
A17503000985_24251251_c2_1868	1016	4788 129 42
Description NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000985_24251400_£2_508	1017	4789 792 263 721 2.9e-71
Description		
gp:[GI:g1854577] [LN:STALYTS] [AC:L42 [DB:genpept-bct1] [DE:Staphylococcus [LE:1849] [RE:2589] [DI:direct]		
ORF Name	NTID	AAID LengthLength score probability
AT7503000985_24251635_±3_1143	1018	4790 123 40
Description		<u> </u>

ORF Name	NTID	AAID	NT AA score probability
AI7503000985_24261062_c1_1458	1019	4791	1773 590 1994 3.7e-206
Description pir:[LN:F70040] [AC:F70040] [PN:sul reductase (ferredoxin)] [OR:Bacillus [LN:BSUB0018] [AC:Z99121:AL009126] [[DB:genpept-bct1] [DE:Bacillus subti 3399551to 3609060.] [NT:similar to s [DI:complement]	subtil GN:yvgQ lis com	is] [I] [FN: plete	unknown] [OR:Bacillus subtilis] genome (section 18 of 21): from
[DI. complement)			
ORF Name	NTID	AAID	NT AA LengthLength score probability
AI7503000985_24269812_f1_414	1020	4792	627 208 285 4.7e-25
Description gp:[GI:e1312907:g3355681] [LN:SC1C2] [GN:SC1C2.14c] [OR:Streptomyces coel coelicolor cosmid 1C2.] [NT:SC1C2.14 [RE:15516] [DI:complement]	icolor]	[DB:g	
ORF Name	NTID	AAID	NT AA score probability
AI7503000985_24273375_c2_1749	1021	4793	
Description gp:[GI:g4574118] [LN:AF009415] [AC:A [OR:Staphylococcus xylosus] [DB:genp transporter (cudT), putativeregulato dehydrogenase(cudA), and choline deh [LE:811] [RE:2433] [DI:direct]	ept-bct ry prot	2] [DE ein (c	:Staphylococcus xylosus choline
ORF Name AI7503000985_24275187_£2_858 Description	NTID 1022	AAID 4794	NT AA score probability [129] 42
NO-HIT	1000		
ORF Name A17503000985_24303775_f2_444	<u>NTID</u>	<u>AAID</u>	NT AA LengthLength score probability
Description NO-HIT			
ORF Name	NTID	AAID	NT AA score probability
AI7503000985 24304187 c2 1728	1024	4796	153 50

Description NO-HIT

ORF Name	NTID	AAID	LengthLength score probability
A17503000985_24304712_£1_311	1025	4797	1674 557 1015 2.1e-102
Description			
sp:[LN:DCIP_ENTCL] [AC:P23234] [GN:I [DE:DECARBOXYLASE)] [SP:P23234] [DB: [PN:indolepyruvate decarboxylase,] [pyrophosphate-binding domain homolog >gp:[GI:d1014947:g216677] [LN:ENTIPD cloacae (strain FERM BP-1529) genomi indolepyruvate decarboxylase.] [NT:i [DI:direct]	swisspr CL:acet y] [OR: C] [AC: c DNA]	ot] >p olacta Entero D90214 [DB:ge	ir:[LN:S16013] [AC:S16013] te synthase large chain:thiamine bacter cloacae] [EC:4.1.1] [DB:pir2] [OR:Enterobacter cloacae] [SR:E. npept-bct1] [DE:E. cloacae gene for
ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000985_24333286_f2_802	1026	4798	126 41
Description			
NO-HIT	MMP (day 1) 1 to an ellipse in any and a second at		
ORF Name	NTID	AAID	NT AA score probability
A17503000985_24337791_c3_2068	1027	4799	1113 370
Description			
NO-HIT			
ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000985_24337807_c3_2076	1028	4800	1890 629 1795 4.6e-185
Description pir: [LN:G70040] [AC:G70040] [PN:sul reductase (NADPH):flavodoxin homolog [OR:Bacillus subtilis] [DB:pir2] >gp [AC:Z99121:AL009126] [GN:yvgR] [FN:u [DE:Bacillus subtilis complete genom [NT:similar to sulfite reductase] [L	y: NADP :[GI:e1: nknown] e (sect:	Hfer: 186032 [OR:Baion 18	:g2635857] [LN:BSUB0018] acillus subtilis] [DB:genpept-bct1] of 21): from 3399551to 3609060.]
ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000985_24338217_c2_1747	1029	4801	597 198 372 2.8e-34
Description			
<pre>gp:[GI:g4098082] [LN:LLU73336] [AC:U activator] [GN:nrdG] [OR:Lactococcus anaerobic ribonucleotide reductase (activator protein (nrdG) genes, comple</pre>	lactis nrdD) an	[DB:q	genpept-bct2] [DE:Lactococcus lactis
ORF Name	NTID	AAID	<u>NT AA</u> LengthLength
AI7503000985_24351562_f3_1230	1030	4802	255 84
Description			
NO-HIT			

ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000985_24353390_c2_1882	1031	4803	696 231 421 1.8e-39
Description pir: [LN:S76993] [AC:S76993] [PN:hyp dehydrogenase:short-chain alcohol de [SR:PCC 6803, , PCC 6803] [SR:PCC 68 [LN:SYCSLRG] [AC:D64005:AB001339] [P [SR:Synechocystis sp. (strain:PCC680 PCC6803 complete genome, 24/27, 3002 [RE:35078] [DI:direct]	hydrogen 803,] [I PN:hypoth 93) DNA]	nase h DB:pir netica [DB:g	nomology] [OR:Synechocystis sp.] [2] >gp:[GI:d1011336:g1001805] [al protein] [OR:Synechocystis sp.] [genpept-bct1] [DE:Synechocystis sp.
ORF Name	NTID	AAID	NT AA score probability
A17503000985_24353392_c2_1954	1032	4804	<u> </u>
Description			
product] [LE:181] [RE:2586] [DI:dire [AC:AF029224:AF029225] [PN:NirB] [GN [DB:genpept-bct2] [DE:Staphylococcus	nce 1 fi ct] >gp: [:nirB] carnos	rom Pa :[GI:g [OR:St ıs nir	tent EP0805205.] [NT:unnamed protein g4433639] [LN:AF029224]
ORF Name	NTID	AAID	NT AA score probability
A17503000985_24406542_f3_1293	1033	4805	339 112 513 3.2e-49
Description gp:[GI:g2914128] [LN:SEU43366] [AC:U epidermidis] [DB:genpept-bct2] [DE:S intercellular adhesion:IcaR, IcaA, I [RE:2268] [DI:direct]	taphylo	coccus	
ORF Name	NTID	AAID	NT AA score probability
A17503000985_24406952_c1_1434	1034	4806	447 148 188 8.9e-15
Description sp:[LN:PETP_RHOCA] [AC:P31078] [GN:P [SR:,RHODOPSEUDOMONAS CAPSULATA] [DE >pir:[LN:S22631] [AC:S22631:S21001] capsulatus] [DB:pir2] >gp:[GI:e49248 [PN:protein of unknown function] [GN [DB:genpept-bct1] [DE:R.capsulatus p operon in front of fbc operon] [SP:P	:PETP PR [PN:pet :g133380 :petP] etP, pet	ROTEIN P pro 02] [Li [OR:Rho R, and	I] [SP:P31078] [DB:swissprot] Dtein] [GN:petP] [OR:Rhodobacter LN:RCPETPR] [AC:Z12113:S42067] Lodobacter capsulatus] Ld fbcF genes.] [NT:part of the petPR
ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000985_24415887_c2_1761	1035	4807	927 308 123 2.7e-05
Description			
<pre>gp:[GI:g3955198] [LN:AF022796] [AC:A carnosus] [DB:genpept-bct2] [DE:Stap biosynthetic genecluster, complete s [LE:109] [RE:894] [DI:direct]</pre>	hylococo	cus ca	

ORF Name	NTID AAID NT AA score probability
A17503000985 24416068 cl 1624	
[OR:Bacillus subtilis] [DB:pir2] >gp [AC:Z99118:AL009126] [GN:ytmI] [FN:u [DE:Bacillus subtilis complete genom [NT:similar to hypothetical proteins [DI:complement] >gp:[GI:e1185812:g26 [FN:unknown] [OR:Bacillus subtilis] genome (section 16 of 21): from 2997 proteins from B. subtilis] [LE:9552] [LN:AF008220] [AC:AF008220] [PN:YtmI [DB:genpept-bct2] [DE:Bacillus subtilis]	nserved hypothetical protein ytmI] [GN:ytmI] p:[GI:e1184188:g2635404] [LN:BSUB0015] unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] ne (section 15 of 21): from 2795131to 3013540.] s from B. subtilis] [LE:212192] [RE:212728] s35423] [LN:BSUB0016] [AC:Z99119:AL009126] [GN:ytmI] [DB:genpept-bct1] [DE:Bacillus subtilis complete 7771to 3213410.] [NT:similar to hypothetical [RE:10088] [DI:complement] >gp:[GI:g2293246] [GN:ytmI] [OR:Bacillus subtilis] tilis rrnB-dnaB genomic region.] [NT:similar to a
hypothetical 19 kD protein from B.]	[LE:170339] [RE:170875] [DI:direct]
ORF Name	NTID AAID LengthLength score probability
A17503000985_24421937_c3_1999	1037 4809 1269 422 778 2.7e-77
<pre>subtilis] [DB:pir2] >gp:[GI:e1182844 [GN:yfhI] [FN:unknown] [OR:Bacillus</pre>	
ORF Name	NTID AAID LengthLength score probability
A17503000985_24429663_c1_1444	1038 4810 933 310 326 2.1e-29
[DE:(EC 2.7.3)] [SP:P42422] [DB:sw [PN:two-component sensor histidine k subtilis] [DB:pir2] >gp:[GI:d1003811 protein] [GN:B65E] [OR:Bacillus subt trpC2)) DNA] [DB:genpept-bct1] [DE:B the iol operon.] [NT:homologous to s [RE:12870] [DI:direct] >gp:[GI:e1184 [GN:yxdK] [FN:unknown] [OR:Bacillus complete genome (section 21 of 21):	XXDK:B65E] [OR:BACILLUS SUBTILIS] [EC:2.7.3] vissprot] >pir:[LN:H70073] [AC:H70073] xinase homolog yxdK] [GN:yxdK] [OR:Bacillus 1:g709992] [LN:BACIOLO] [AC:D14399] [PN:hypothetical xilis] [SR:Bacillus subtilis (strain:BGSC 1A1 (168 Bacillus subtilis 15 kb chromosome segment contains xensor protein BvgC, His protein] [LE:11893] x690:g2636511] [LN:BSUB0021] [AC:Z99124:AL009126] xubtilis] [DB:genpept-bct1] [DE:Bacillus subtilis xfrom 3999281to 4214814.] [NT:similar to xe] [SP:P42422] [LE:70819] [RE:71796]
ORF Name	NTID AAID NT AA score probability
[OR:Lactococcus lactis] [SR:Lactococ	PN:alpha-acetolactate synthase] [GN:als] cus lactis (strain DSM 20384, sub_species lactis) rus lactis alpha-acetolactate synthase (als) gene

DNA] [DB:genpept-bct1] [DE:Lactococcus lactis alpha-acetolactate synthase (als) gene, completecds.] [LE:1232] [RE:2896] [DI:direct] >gp:[GI:g809618] [LN:A23961] [AC:A23961] [PN:alpha-acetolactate synthase] [OR:Lactococcus lactis] [DB:genpept-pat] [DE:L. lactis alpha-acetolactate synthase gene.] [LE:550] [RE:2214] [DI:direct]

ORF Name	NTID	AAID	NT LengthL	ength score	probability
A17503000985_24500300_c1_1462	1040	4812	7 1206	101 1208	7.3e-123
Description			<i></i>		
pir:[LN:B69877] [AC:B69877] [PN:sulfile] [CL:Synechocystis sulfate adenylylta [OR:Bacillus subtilis] [DB:pir2] >gg [PN:putative sulfate adenylyltransfe [DB:genpept-bct1] [DE:Bacillus subtilis] [DI:direct] >gp:[GI:e1185151:g263393] [FN:unknown] [OR:Bacillus subtilis] genome (section 9 of 21): from 15984 adenylyltransferase] [LE:33187] [RE:	cansfera o:[GI:e3 erase] [ilis pyr B2] [LN: [DB:gen 121to 18	se: su 32185: GN:yln E to y BSUB00 pept-b 07200.	lfate ad g2462958 B] [OR:E loA gene 09] [AC: ctl] [DE] [NT:si	denylyltrans [] [LN:BSPYR Bacillus sub region.] [Z99112:AL00 C:Bacillus s	sferase homology] REYLO] [AC:AJ000974] Otilis] [LE:2374] [RE:3522] O9126] [GN:ylnB] Subtilis complete
ORF Name	NTID	AAID	<u>NT</u> LengthL	AA score	probability
AI7503000985_245443_c1_1548	1041	4813	696 2	231 375	1.4e-34
Description pir: [LN:F69879] [AC:F69879] [PN:pho pir: [LN:F69879] [AC:F69879] [DB:pir2] > [OR:Bacillus subtilis] [DB:pir2] > [AC:Z99112:AL009126] [GN:yloW] [FN:u [DE:Bacillus subtilis complete genom [NT:similar to phosphoglycerate dehy pogp: [GI:e323511:g2337814] [LN:BSY139 [GN:yloW] [FN:unknown] [OR:Bacillus genomic DNA from the spoVM region.]	egp:[GI: unknown] ne (sect vdrogena 037] [AC subtili	e11851 [OR:B ion 9 se] [L :Y1393 s] [DB	76:g2633 acillus of 21): E:59221] 7] [PN:p :genpept	subtilis] [from 159842 [RE:59883] sutative Yha -bct1] [DE:	DB:genpept-bct1] The left of t
ORF Name	NTID	AAID	<u>NT</u> LengthL	AA ength	probability
AI7503000985_24611567_c2_1831	1042	4814			1.7e-45
Description pir:[LN:S36209] [AC:S36209] [PN:depenthracis] [DB:pir2] >gp:[GI:d100363] [GN:dep] [OR:Bacillus anthracis] [SR DNA] [DB:genpept-bct1] [DE:Bacillus cds.] [LE:252] [RE:1652] [DI:direct]	2:g4360 Bacillanthrac	34] [L us ant	N:BACDEP hracis (] [AC:D1403 strain:Davi	7] [PN:ORF] s) plasmid:pTE702
ORF Name	NTID	AAID	NT LengthL		probability

Description NO-HIT

ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	score	probability
AI7503000985_24617262_c1_1501	1044	4816		274		5.7e-68
Description sp:[LN:PANB_BACSU] [AC:P52996] [GN:Pi [DE:(KETOPANTOATE HYDROXYMETHYLTRANS] >pir:[LN:G69671] [AC:G69671] [PN:ket] [CL:3-methyl-2-oxobutanoate hydroxy [DB:pir2] >gp:[GI:g1146240] [LN:BACYI hydroxymethyltransferase] [GN:panB] subtilis] [DB:genpept-bct1] [EC:2.1.2 genes, qcrABC genes,ypjABCDEFGHI gene gene,aspB gene, asnS gene, dnaD gene of identity to the 3-methyl-2-oxobuta >gp:[GI:e1183688:g2634661] [LN:BSUBOO hydroxymethyltransferase] [GN:panB] subtilis] [DB:genpept-bct1] [EC:2.1.2 (section 12 of 21): from 2195541to 24 [DI:complement]	FERASE) topanto ymethyl PIA] [A [FN:pan 2.11] [i es, bir , nth g anoate] 012] [A [FN:pan 2.11] [i] [SP:1 ate hyd transfe C:L4770 tothen: DE:Bac: A gene ene and [LE:1: C:Z991: tothena	p52996] droxyme erase] p9] [PN ic acid illus s panBC d ypoC s 3293] [15:AL00 ate bio illus s	[DB:sthyltr [OR:Ba:ketop biosy ubtili D gene gene, RE:141 9126] synthe	wisspansfer cillus antoat nthes: s (close, direction complete) [PN:kessis] s comp	cot] case panB] [GN:panB s subtilis] ce is] [OR:Bacillus one YAC15-6B) ypiABF nG gene, ypmB etecds's.] [NT:47.1% DI:direct] etopantoate [OR:Bacillus olete genome
ORF Name A17503000985_24640910_c2_1654 Description NO-HIT	NTID 1045	AAID 4817	NT Length]	AA Length 172	score	probability
ORF Name A17503000985_24641932_c2_1734 Description gp:[GI:g4104595] [LN:AF036964] [AC:AN [OR:Lactobacillus sakei] [DB:genpeptregulator (rrpl) and putative histiding member of a two-component regulatory	-bct2] ne kina:	4818 PN:p DE:Lac	outative ctobaci	e resp llus s	onse make pu	tative response c cds.] [NT:Rrp1;
ORF Name AI7503000985_24643930_c3_2138 Description sp:[LN:YOHK_ECOLI] [AC:P33373] [GN:YOPROTEIN IN PBPG-CDD INTERGENIC REGION [AC:E64982] [PN:yohK protein] [GN:yoPROTEIN] [SP:QDB:pir2] >gp:[GI:g1788464] [LN:AE000Ctransporter] [GN:yohK] [FN:putative to [DB:genpept-bct2] [DE:Escherichia colecompletegenome.] [NT:o231; residues 1] [DI:direct]	N] [SP:1 bhK] [0 0303] [<i>i</i> cranspor Li K-12	R:ESCHE P33373] CL:yohk AC:AE00 ct; Not MG1655	ERICHIA [DB:st [prote: 0303:U0 class: sections	COLI] wissprain] [O 00096] ified] on 193	[DE:Hot] >pR:Esch [PN:p [OR:E	oir:[LN:E64982] Perichia coli] Putative seritonin Packscherichia coli] Poof the
ORF Name AI7503000985_24646963_c3_2072 Description gp:[GI:e315090:g2791905] [LN:SSK3MECF sciuri] [DB:genpept-bct1] [DE:S.sciur		4820 :Y13052] [GN:0	RF454	[OR:	

		NT AA
ORF Name	NTID	AAID LengthLength score probability
AI7503000985_24648377_c3_2261	1049	4821 1146 381 1301 1.0e-132
Description		
[OR:Staphylococcus carnosus] [DB:ger [LE:538] [RE:1704] [DI:direct] >gp: transporter] [GN:narT] [OR:Staphyloc	npept-pa [GI:g252 coccus c :Staphyl	A67169] [PN:NART GENE] [FN:NITRATE TRANSPORT] At] [DE:Sequence 9 from Patent EP0805205.] Ap402] [LN:SCU40014] [AC:U40014] [PN:nitrate carnosus] [SR:Staphylococcus carnosus Accoccus carnosus nitrate transporter (narT) Ap56] [DI:direct]
ORF Name	NTID	AAID NT AA score probability
A17503000985_24648502_c3_2088	1050	4822 468 155 228 5.1e-19
KD PROTEIN IN ECSC-PBPF INTERGENIC F [AC:B40614:F69832] [PN:conserved by (pbpF 5' region)] [GN:yhgC] [OR:Bacillu DNA] [DB:genpept-bct1] [DE:Bacillus end.] [NT:product unknown] [LE:247] [LN:BSUB0006] [AC:Z99109:AL009126] [DB:genpept-bct1] [DE:Bacillus subtito1209940.] [NT:alternate gene name: [LE:83202] [RE:83702] [DI:complement [AC:Y14083] [PN:Hypothetical protein [DB:genpept-bct1] [DE:Bacillus subtito1209940.] [NT:Hypothetical protein [DB:genpept-bct1] [DE:Bacillus subtito1209940.]	REGION] ypotheti cillus s us subti subtili [RE:747 [GN:yhgC ilis com : yixC; t] >gp:[n] [GN:y ilis chr	lis] [SR:Bacillus subtilis (strain W168) s penicillin-binding protein (pbpF) gene, 5'] [DI:complement] >gp:[GI:e1183012:g2633346]] [FN:unknown] [OR:Bacillus subtilis] splete genome (section 6 of 21): from 999501 similar to hypothetical] [SP:P38049] GI:e325006:g2226228] [LN:BSY14083] rixC] [OR:Bacillus subtilis]
ORF Name	NTID	AAID NT AA score probability
A17503000985_24648551_f2_791 Description	1051	4823 174 57 108 6.6e-06
[LE:394] [RE:1083] [DI:complement] > [OR:Staphylococcus aureus] [SR:Staphylococcus [DB:genpept-bct1] [DE:Staphylococcus	emolytic 272 ORF1 egp:[GI: nylococc aureus	us strain=Y176] [DB:genpept-bct1] and ORF2 genes, completecds.] [NT:ORF2] g295162] [LN:STAMECRA] [AC:L14017]
ORF Name	NTID	AAID NT AA score probability
A17503000985_24652312_f1_419	1052	4824 1617 538 1137 2.4e-115
Description [72]		
gp:[G1:g4835822] [LN:AF102174] [AC:A	1102174] [PN:glycine betaine transporter BetL]

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[GN:betL] [OR:Listeria monocytogenes] [DB:genpept-bct2] [DE:Listeria monocytogenes glycine betaine transporter BetL (betL)gene, complete cds.] [LE:209] [RE:1732]

[DI:direct]

ORF Name	NTID AAID NT AA score probability
A17503000985_24665957_c3_2149	1053 4825 1494 497 1335 2.5e-136
[DE:XYLULOSE KINASE, (XYLULOKINA [AC:D69735] [PN:xylulose kinase subtilis] [DB:pir2] >gp:[GI:g175 [GN:xylB] [OR:Bacillus subtilis] YnbA (ynbA), YnbB (ynbB), GlnR(g (ynaB), YnaC(ynaC), YnaD (ynaD), YnaI (ynaI), YnaJ (ynaJ), xylan xylose isomerase (xylA), xylulos and YncE (yncE)genes, complete copp:[GI:e1183420:g2634145] [LN:B. [GN:xylB] [FN:xylose metabolism] [DE:Bacillus subtilis complete genes.]	GN:XYLB] [OR:BACILLUS SUBTILIS] [EC:2.7.1.17] ASE)] [SP:P39211] [DB:swissprot] >pir:[LN:D69735] ExylB] [GN:xylB] [CL:xylulokinase] [OR:Bacillus 10125] [LN:BSU66480] [AC:U66480] [PN:xylulose kinase] [DB:genpept-bctl] [DE:Bacillus subtilis SpoVK (spoVK), rlnR), glutamine synthetase (glnA), YnaA (ynaA), YnaB YnaE (ynaE), YnaF (ynaF), YnaG (ynaG), YnaH(ynaH), beta-1,4-xylosidase (xynB),xylose repressor (xylR), se kinase(xylB), YncB (yncB), YncC (yncC), YncD (yncD) 2ds.] [LE:19399] [RE:20898] [DI:direct] 2SUB0010] [AC:Z99113:AL009126] [PN:xylulose kinase] [OR:Bacillus subtilis] [DB:genpept-bct1] [EC:2.7.1.17] 2enome (section 10 of 21): from 1781201to 2014980.] 3SP:P39211] [LE:111450] [RE:112949] [DI:direct]
ORF Name AI7503000985_24720257_f1_131 Description NO-HIT	NTID AAID NT AA score probability 1054 4826 138 45
[GN:ywtB] [OR:Bacillus subtilis [AC:Z99122:AL009126] [GN:ywtB] [EDE:Bacillus subtilis complete go [NT:similar to capsular polyglut [DI:complement] >gp:[GI:e308090:gor:Bacillus subtilis] [DB:genpe] genes.] [NT:product highly similated [DI:direct] >gp:[GI:e1184494:g265] [FN:unknown] [OR:Bacillus subtilis]	NTID AAID LengthLength Score probability [1055] 4827 [1101] 366 [353] [2.9e-32] [scapsular polyglutamate biosynthesis homolog ywtB] [stapsular polyglutamate biosynthesis homolog ywtB] [stapsular polyglutamate biosynthesis homolog ywtB] [stapsular polyglutamate biosynthesis [LN:BSUB0019] [stapsular polyglutamate biosynthesis] [DB:genpept-bct1] [stapsular polyglutamate subtilis] [DB:genpept-bct1] [stapsular polyglutamate biosynthesis] [LE:99788] [RE:100930] [g1894767] [LN:BSZ92954] [AC:Z92954] [GN:ywtB] [stapsular polyglutamate polyglutamate ganome to 3809700.] [NT:similar to capsular polyglutamate ganome to 3809700.] [NT:similar to capsular polyglutamate ganome to 3809700.] [DI:complement]
[OR:Lactobacillus sakei] [DB:genpartial cds; LaaB (laaB),putative	NTID AAID NT AA LengthLength score probability 1056 4828 420 139 374 1.7e-34 AC:AF115391] [PN:ribose permease RbsD] [GN:rbsD] pept-bct2] [DE:Lactobacillus sakei LaaA (laaA) gene, e acetate kinase AckA (ackA), LaaC (laaC) genes, e sequence; and LaaE (laaE) gene, partialcds.]
-	NTID AAID NT AA Score probability 1057 4829 900 299 145 7.3e-08 OR:PSEUDOMONAS FLUORESCENS] [EC:3.1.1.2] DROLASE)] [SP:P22862] [DB:swissprot]

ORF Name	NTID	AAID	NT AA	probability
			LengthLength	<u> </u>
A17503000985_24797900_c1_1361	1058	4830	213 70	
<u>Description</u>				
NO-HIT				
ORF Name	NTID	AAID	NT AA LengthLength score	probability
A17503000985_24812502_c1_1521	1059	4831	126 41	
Description				
NO-HIT				
ORF Name	NTID	AAID	NT AA score	probability
A17503000985_24814838_c2_1789	1060	4832		.1e-32
Description			<u> </u>	
pir:[LN:S76790] [AC:S76790] [PN:hyp [OR:Synechocystis sp.] [SR:PCC 6803, >gp:[GI:d1019435:g1653791] [LN:D9091 [OR:Synechocystis sp.] [SR:Synechocy [DE:Synechocystis sp. PCC6803 comple [NT:ORF_ID:slr1563] [LE:99350] [RE:1	, PCC (6] [AC:1 stis sp te genor	6803] D90916 . (str me, 26	[SR:PCC 6803,] [DB: :AB001339] [PN:hypot ain:PCC6803) DNA] [D /27, 3270710-3418851	pir2] hetical protein] B:genpept-bct1]
ORF Name	NTID	AAID	NT AA LengthLength	probability
A17503000985_24817202_c1_1512	1061	4833	1809 602 799 2	.3e-116
Description		L		
pir:[LN:C69975] [AC:C69975] [PN:acy subtilis] [DB:pir2] >gp:[GI:g1934616] protein YrhL] [GN:yrhL] [OR:Bacillus cysteine synthase (yrhA), cystathion formate dehydrogenasechain A (yrhE), YrhH(yrhH), regulatory protein (yrhI hypothetical protein YrhL (yrhL), pu factor SigV (sigV) and YrhO (yrhO)ge:cds.] [NT:similar to Haemophilus inf [DI:complement] >gp:[GI:e1183944:g26 [FN:unknown] [OR:Bacillus subtilis] genome (section 14 of 21): from 2599 [LE:171138] [RE:173042] [DI:direct]	[LN:Bs subtil: inegamma YrhF (y), cytootative anes, corluenzae 35160]	SU9387 is] [D a-lyas yrhF), chrome anti-S mplete hypot [LN:BS pept-b	4] [AC:U93874] [PN:h B:genpept-bct1] [DE: e (yrhB), YrhC (yrhC formate dehydrogena P450 102 (yrhJ), Yr igV factor(yrhM), RN cds, and YrhP (yrhP hetical] [LE:13904] UB0014] [AC:Z99117:A ct1] [DE:Bacillus su	ypothetical Bacillus subtilis (), YrhD (yrhD), se (yrhG), hK(yrhK), A polymerase sigma () gene, partial [RE:15808] L009126] [GN:yrhL] btilis complete
ORF Name	NTID	AAID	NT AA LengthLength score	probability
A17503000985_24855337_c2_1957	1062	4834		.8e-101
Description		•		
<pre>gp:[GI:e1429602:g4756154] [LN:A67161 carnosus] [DB:genpept-pat] [DE:Seque: product] [LE:9942] [RE:10625] [DI:di: [AC:AF029224:AF029225] [PN:NarI] [GN [DB:genpept-bct2] [DE:Staphylococcus</pre>	nce 1 fi rect] >@ :narI]	com Pa gp:[GI [OR:St	tent EP0805205.] [NT :g3929525] [LN:AF029 aphylococcus carnosu	unnamed protein 224] s]
[NT:similar to Escherichia coli nitra			——————————————————————————————————————	·

[DI:direct]

ORF Name	NTID AAID NT AA score probability
AI7503000985_250178_c3_2162	1063 4835 234 77 82 0.0015
Description	
[OR:Mycobacterium tuberculosis] [DB: [AC:Z95554:AL123456] [PN:hypothetica tuberculosis] [DB:genpept-bct1] [DE:	oothetical protein Rv1615] [GN:Rv1615] pir2] >gp:[GI:e316881:g2113899] [LN:MTCY01B2] al protein Rv1615] [GN:Rv1615] [OR:Mycobacterium Mycobacterium tuberculosis H37Rv complete genome; B2.07), len: 146. Function: unknown] [LE:7451]
ORF Name	NTID AAID NT AA score probability
A17503000985_2507950_f1_199	1064 4836 216 71
Description NO-HIT	
ORF Name	NTID AAID Install score probability
A17503000985 251 £1 433	LengthLength
Description	
NO-HIT	
ORF Name	NTID AAID NT AA score probability
A17503000985_25398426_f1_211	1066 4838 198 65
Description NO-HIT	
ORF Name	NTID AAID NT AA probability
AI7503000985_25413577_f3_1032	1067 4839 129 42
Description	
NO-HIT	
ORF Name	NTID AAID NT AA score probability
A17503000985_25429700_c2_1683	1068 4840 1356 451 158 1.8e-08
Description	
[DB:genpept-bct1] [DE:B.cereus DNA f	8] [AC:Y11138] [GN:ORF1] [OR:Bacillus cereus] or ORF1, ORF2 and ORF3 (2402 bp).] [NT:shows weak ORF] [LE:156] [RE:1373] [DI:direct]
ORF Name	NTID AAID NT AA probability
A17503000985_25431558_f1_63	1069 4841 705 234 855 1.9e-85
Description	
aureus] [SR:Staphylococcus aureus st	
	protein LrgA (lrgA) and LrgB(lrgB) genes, complete lar to E.coli vohK [LE:805] [RE:1506] [DI:direct]

ORF Name	NTID	AAID NT AA probability LengthLength score probability
A17503000985_25433452_f1_61	1070	4842 1791 596 1867 1.1e-192
Description		
<pre>gp:[GI:g862312] [LN:STALYTS] [AC:L42 [DB:genpept-bct1] [DE:Staphylococcus [RE:1846] [DI:direct]</pre>		N:lytS] [OR:Staphylococcus aureus] lytS and lytR genes, complete cds.] [LE:92]
ORF Name	NTID	AAID <u>NT AA</u> <u>score</u> <u>probability</u>
A17503000985_25570262_f1_358	1071	4843 144 47
Description		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000985_25578215_c2_1740	1072	4844 543 180 483 4.9e-46
Description		
<pre>gp:[GI:e316582:g2791909] [LN:SSK11ME sciuri] [DB:genpept-bct1] [DE:S.sciu [RE:4914] [DI:direct]</pre>		:Y13094] [GN:ORF141] [OR:Staphylococcus gene, strain K11 (792).] [LE:4489]
ORF Name	NTID	AAID NT AA score probability LengthLength
A17503000985_2557962_t2_698	1073	4845 1377 458 953 7.7e-96
Description		
FAECALIS] [EC:1.6.99.3] [DE:NADH OXI >pir:[LN:S26965] [AC:S26965] [PN:NA faecalis] [DB:pir2] >gp:[GI:g47045]	DASE, (1 DH oxida [LN:SFNO [DB:geng	ASE] [CL:NADH peroxidase] [OR:Enterococcus DXAA] [AC:X68847:S45681] [PN:NADH oxidase] Dept-bct1] [DE:S.faecalis nox gene for NADH
ORF Name	NTID	AAID NT AA score probability LengthLength
AI7503000985_25580425_f1_74	1074	4846 135 44
Description	<u> </u>	
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
AI7503000985_25585932_c1_1598	1075	4847 276 91 254 9.0e-22
Description		
<pre>gp:[GI:d1039105:g4514322] [LN:AB0133 halodurans] [SR:Bacillus halodurans halodurans C-125 ydeI gene, complete</pre>	(strain:	C-125) DNA] [DB:genpept-bct1] [DE:Bacillus

ORF Name	NTID	AAID	NT AA score probability
A17503000985 25600015 c3 2090	1076	4848	Length Length
Description	1070	1010	1032 343 1301 2.96-100
pir:[LN:E69989] [AC:E69989] [PN:ace ligase homology] [OR:Bacillus subtil >gp:[GI:e1185829:g2635440] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-b 16 of 21): from 2997771to 3213410.] [RE:26574] [DI:complement] >gp:[GI:g [GN:ytcI] [OR:Bacillus subtilis] [DB	is] [EC 016] [AG ct1] [DI [NT:sim: 2293232] :genpept	:6.2.1 C:Z991 E:Baci ilar t] [LN: t-bct2	119:AL009126] [GN:ytcI] [FN:unknown] illus subtilis complete genome (section to acetate-CoA ligase] [LE:24979] :AF008220] [AC:AF008220] [PN:YtcI]
ORF Name	NTID	AAID	NT AA LengthLength score probability
AI7503000985_25626875_t2_653	1077	4849	147 48
Description			
NO-HIT			
ORF Name	NTID	AAID	NT AA LengthLength score probability
AI7503000985_25666427_£3_1335	1078	4850	306 101 84 0.0061
Description		<u>-</u>	
SULFONATES BINDING PROTEIN PRECURSOR [AC:I39927:C69817] [PN:ABC transport [OR:Bacillus subtilis] [DB:pir2] >gp [OR:Bacillus subtilis] [SR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis] [NT:Likely N-terminal signal sequence >gp:[GI:e1182873:g2633207] [LN:BSUB06] [OR:Bacillus subtilis] [DB:genpept-bc5 of 21): from 802821 to1011250.] [NT:DB:genpept-bc5 of 21): from 802821 to1011250.]	[SP:P4] ter (bir :[GI:g43] subtilis lis orff e, follo 005] [AC ct1] [DE T:alterr [DI:dire lipopro btilis y	40400] nding 38472] s (ind K, orf owed b C:Z991 E:Baci nate g ect] > otein j	lipoprotein) homolog ygbA] [GN:ygbA] [LN:BACORFKLM] [AC:L16808] dividual_isolate MS11) (library: Tn91] fL and orfM, complete cds's.] by] [LE:69] [RE:1067] [DI:direct] 108:AL009126] [GN:ygbA] [FN:unknown] illus subtilis complete genome (section gene name: yzeA; similar to ABC] >gp:[GI:e308630:g1903039] [LN:BSZ93102] precursor] [GN:yzeA] [OR:Bacillus ,M,N,O,P,Q,R,S,T], yzdB and yze[A,C}
ORF Name	NTID	AAID	NT AA score probability
AI7503000985_25961087_f1_123	1079	4851	
Description			
KD PROTEIN IN GLVBC 3'REGION] [SP:P54 [PN:conserved hypothetical protein yipg:[GI:e1182814:g2633148] [LN:BSUB00	4721] [EfiE] [GN 005] [AC ct1] [DE T:simila :95553] [GN:yfi aplotype	DB:swis N:yfiE C:Z9910 E:Bacil ar to l [DI:d: [DI:d: e:haplo	illus subtilis complete genome (section hypothetical proteins from B. direct] >gp:[GI:d1009744:g1486247] FN:unknown] [OR:Bacillus subtilis] loid) DNA] [DB:genpept-bct1]

[DI:direct]

ORF Name AI7503000985_25970952_f1_316 Description NO-HIT	NTID	AAID NT AA score probability 4852 141 46
ORF Name AI7503000985_26017278_c1_1441 Description NO-HIT	NTID 1081	AAID NT AA score probability LengthLength 4853 132 43
[SR:PCC 6803, , PCC 6803] [SR:PCC 68 [LN:SYCSLLE] [AC:D64003:AB001339] [F [SR:Synechocystis sp. (strain:PCC680	03,] [1 N:hypot] 3) DNA]	AAID NT AA LengthLength score probability 4854 1818 605 522 3.6e-50 al protein sll0556] [OR:Synechocystis sp.] DB:pir2] >gp:[GI:d1011128:g1001236] hetical protein] [OR:Synechocystis sp.] [DB:genpept-bct1] [DE:Synechocystis sp. 8766.] [NT:ORF_ID:sll0556] [LE:45438]
opuBC] [GN:opuBC] [OR:Bacillus subti [LN:BSUB0018] [AC:Z99121:AL009126] [[GN:opuBC] [FN:high affinity transpot [DB:genpept-bct1] [DE:Bacillus subti 3399551to 3609060.] [NT:alternate ge [DI:complement] >gp:[GI:g2293449] [I protein precursor] [GN:opuBC] [OR:Ba subtilis choline transport system in (opuBB), choline binding proteinprec	PN:chol: ort of cl lis comp ne name N:AF008 cillus s cluding ursor (c	ine ABC transporter (choline-binding) holine] [OR:Bacillus subtilis] plete genome (section 18 of 21): from : proX] [LE:60016] [RE:60936] 930] [AC:AF008930] [PN:choline binding subtilis] [DB:genpept-bct2] [DE:Bacillus ATPase(opuBA), transmembrane protein opuBC) and transmembrane protein (opuBD) part of choline uptake system; OpuBC;
ORF Name A17503000985_26207537_f3_1201 Description NO-HIT	NTID 1084	AAID NT AA score probability LengthLength 4856 129 42
ORF Name AI7503000985_26213885_c3_2215 Description NO-HIT	NTID	AAID NT AA score probability 4857 132 43

			>107	7.7		
ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	score	probability
A17503000985_26220077_t2_445	1086	4858	447	148	259	2.7e-22
Description		<u> </u>		•	·	
pir: [LN:E69857] [AC:E69857] [PN:con [OR:Bacillus subtilis] [DB:pir2] >gp [PN:YkmA] [GN:ykmA] [FN:unknown] [OR subtilis 168 56 kb DNA fragment betw [DI:complement] >gp:[GI:e1183335:g26 [FN:unknown] [OR:Bacillus subtilis] genome (section 7 of 21): from 11943 proteins] [LE:186508] [RE:186951] [D	e:[GI:el e:Bacill een xly 33669] [DB:gen 91to 14	181515 us sub A and [LN:BS pept-b 11140.	:g26320 tilis] ykoR.] UB0007] ct1] [D	35] [I [DB:ge [LE:33 [AC:Z	N:BSAJ: npept-l 559] [1 99110:2	2571] [AC:AJ002571] cct1] [DE:Bacillus RE:34002] AL009126] [GN:ykmA] abtilis complete
ORF Name	NTID	AAID	NT Length1	<u>AA</u> Length	score	probability
AI7503000985_26353411_c1_1391	1087	4859		191		
Description						
NO-HIT			A -			
ORF Name	NTID	AAID	<u>NT</u> LengthI	<u>AA</u> Length	score	probability
AI7503000985_26380265_c3_2042	1088	4860	1263	420	1291	2e-131
Description						
<pre>gp:[GI:e1299582:g3687416] [LN:BLY175 [OR:Bacillus licheniformis] [DB:genp arcA, arcB, arcC and arcD genes.] [L</pre>	ept-bct	1] [EC	:3.5.3.	6] [DE	:Bacill	
ORF Name	NTID	AAID	<u>NT</u> LengthI	<u>AA</u> Length	score	probability
A17503000985_26383512_c1_1506	1089	4861	1398	465	1195 1	7e-121
sp:[LN:YDGF_BACSU] [AC:P96704] [GN:YTRANSPORT PROTEIN IN EXPZ-DINB INTER >pir:[LN:H69782] [AC:H69782] [PN:am [GN:ydgF] [CL:arginine permease] [O >gp:[GI:d1020148:g1881368] [LN:AB001 [SR:Bacillus subtilis (strain:168) Disequence, 148 kb sequence of the reg ACID TRANSPORT PERMIASE.] [LE:139917 >gp:[GI:e1182528:g2632862] [LN:BSUB0	GENIC R ino aci R:Bacil 488] [A NA] [DB ionbetw] [RE:1	EGION] d ABC lus sul C:AB00: :genpe een 35 41293]	[SP:P96 transpon btilis] 1488] [6 pt-bct1] and 47 [DI:cor	6704] rter ([DB:p GN:ydg] [DE: degre mpleme	[DB:swi permeas ir2] F] [OR: Bacillu e.] [N]	Bacillus subtilis] subtilis genome T:PROBABLE AMINO
[OR:Bacillus subtilis] [DB:genpept-b 3 of 21): from 402751 to611850.] [NT [SP:P96704] [LE:203663] [RE:205039] [LN:BSUB0004] [AC:Z99107:AL009126] [DB:genpept-bct1] [DE:Bacillus subtilises]	ct1] [D :simila [DI:com GN:ydgF	E:Bacil r to an plement] [FN:1	llus sub mino aci t] >gp: unknown]	btilis id ABC [GI:e1] [OR:	comple transr 182541: Bacillu	ete genome (section corter (permease)] g2632875] as subtilis]

to813890.] [NT:similar to amino acid ABC transporter (permease)] [SP:P96704] [LE:5713]

[RE:7089] [DI:complement]

ORF Name	NTID	AAID Length Length score probability
A17503000985_26385928_£2_900	1090	4862 1437 478 1441 1.5e-147
Description		
[GN:yfls] [CL:2-oxoglutarate/malate >gp:[GI:e1182747:g2633081] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-k 5 of 21): from 802821 to1011250.] [N	e transle 0005] [A oct1] [D VT:simila gp:[GI:d C:Bacilla lis 35.	7 kb genomic DNA, 70-73 degree
ORF Name	NTID	AAID NT AA score probability
AI7503000985_26429800_f2_605	1091	4863 126 41
Description		
NO-HIT		
(*)		NTII 7.7
ORF Name	NTID	AAID LengthLength score probability
AI7503000985_26449187_c1_1436	1092	4864 129 42
Description		
NO-HIT		
		NT AA
ORF Name	NTID	AAID LengthLength score probability
A17503000985_26571937_f2_570	1093	4865 789 262 838 1.2e-83
Description		
pir:[LN:D69845] [AC:D69845] [PN:thi		
[CL:thiamine biosynthesis protein th		
>gp:[GI:e1183189:g2633523] [LN:BSUBO		C:Z99110:AL009126] [GN:yjbT] [FN:unknown] E:Bacillus subtilis complete genome (section
		lar to thiamin biosynthesis] [LE:50122]
[RE:50892] [DI:direct]		
		NITT A A
ORF Name	NTID	AAID LengthLength score probability
A17503000985 26595641 f2 572	1094	4866
Description		
sp:[LN:APL LACLA] [AC:Q48630] [GN:AF	Ll [OR:	LACTOCOCCUS LACTIS1
[SR:, SUBSPLACTIS:STREPTOCOCCUS LACTI		
- - -		[AC:S39339] [PN:alkaline phosphatase-like
	_	yngC] [OR:Lactococcus lactis] [DB:pir2]
		[PN:alkaline phosphatase like protein] t-bct1] [DE:L.lactis (MG1363) apl gene for

alkaline phosphatase like protein.] [SP:Q48630] [LE:339] [RE:1067] [DI:direct]

pir:[LN:F69653] [AC:F69653] [PN:transcription regulator lrpC] [GN:lrpC] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1182391:g2632725] [LN:BSUB0003] [AC:Z99106:AL009126] [PN:transcriptional regulator (Lrp/AsnC family)] [GN:lrpC] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 3 of 21): from 402751 to611850.] [NT:alternate gene name: ydaI] [LE:72862] [RE:73296] [DI:direct]

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ORF Name NTID AAID Length Length score probability

Description

pir:[LN:E69581] [AC:E69581] [PN:acetoin dehydrogenase E1 component (TPP-dependent beta subunit) acoB] [GN:acoB] [CL:pyruvate dehydrogenase (lipoamide) beta chain] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1182797:g2633131] [LN:BSUB0005] [AC:Z99108:AL009126] [PN:acetoin dehydrogenase El component] [GN:acoB] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 5 of 21): from 802821 to1011250.] [NT:alternate gene name: yfjJ] [LE:76693] [RE:77721] [DI:direct] >gp:[GI:d1025207:g2780394] [LN:D78509] [AC:D78509] [PN:YfjJ] [OR:Bacillus subtilis] [SR:Bacillus subtilis (strain:AC327) DNA] [DB:genpept-bct1] [DE:Bacillus subtilis YfjG-YfjR genes, complete cds.] [LE:8604] [RE:9632] [DI:complement] >gp:[GI:g2245638] [LN:AF006075] [AC:AF006075] [PN:TPP-dependent acetoin dehydrogenase, E1] [GN:acoB] [OR:Bacillus subtilis] [DB:qenpept-bct2] [DE:Bacillus subtilis acetoin dehydrogenase enzyme system gene cluster, ribosomal protein L6-like protein gene, partial cds, TPP-dependentacetoin dehydrogenase, E1 alpha-subunit (acoA), TPP-dependentacetoin dehydrogenase, E1 beta-subunit (acoB), dihydrolipoamideacetyltransferase (acoC) and dihydrolipoamide dehydrogenase (acoL)genes, complete cds, and regulatory protein (acoR) gene, partialcds.] subunit of the El component of the acetoin] [LE:1830] [RE:2858] [DI:direct] >qp:[GI:q2245638] [LN:AF006075] [AC:AF006075] [PN:TPP-dependent acetoin dehydrogenase. E1] [GN:acoB] [OR:Bacillus subtilis] [DB:genpept] [DE:Bacillus subtilis acetoin dehydrogenase enzyme system gene cluster, ribosomal protein L6-like protein gene, partial cds, TPP-dependentacetoin dehydrogenase, E1 alpha-subunit (acoA), TPP-dependentacetoin dehydrogenase, E1 beta-subunit (acoB), dihydrolipoamideacetyltransferase (acoC) and dihydrolipoamide dehydrogenase (acoL)genes, complete cds, and regulatory protein (acoR) gene, partialcds.] subunit of the E1 component of the acetoin] [LE:1830] [RE:2858] [DI:direct]

 ORF Name
 NTID
 AAID
 NT AA Length Length
 score
 probability

 A17503000985_26751887_c1_1608
 1100
 4872
 954
 317
 297
 2.5e-26

Description

sp:[LN:APBA_AQUAE] [AC:O67619] [GN:APBA:AQ_1727] [OR:AQUIFEX AEOLICUS] [EC:1.1.1.169] [DE:REDUCTASE) (KPA REDUCTASE)] [SP:O67619] [DB:swissprot] >pir:[LN:A70449] [AC:A70449] [PN:hypothetical protein aq_1727] [GN:aq_1727] [OR:Aquifex aeolicus] [DB:pir2] >gp:[GI:g2984043] [LN:AE000753] [AC:AE000753:AE000657] [PN:putative protein] [GN:aq_1727] [OR:Aquifex aeolicus] [DB:genpept-bct2] [DE:Aquifex aeolicus section 85 of 109 of the complete genome.] [LE:5968] [RE:6900] [DI:complement]

 ORF Name
 NTID
 AAID
 NT AA Length Length
 score
 probability

 A17503000985_273452_c2_1815
 1101
 4873
 1278
 425
 800
 1.2e-79

Description

sp:[LN:HMDH_ARCFU] [AC:028538] [GN:HMGA:AF1736] [OR:ARCHAEOGLOBUS FULGIDUS]
[EC:1.1.1.34] [DE:REDUCTASE)] [SP:028538] [DB:swissprot] >pir:[LN:G69466] [AC:G69466]
[PN:3-hydroxy-3-methylglutaryl-coenzyme A reductase (mvaA) homolog] [OR:Archaeoglobus fulgidus] [DB:pir2] >gp:[GI:g2648815] [LN:AE000983] [AC:AE000983:AE000782]
[PN:3-hydroxy-3-methylglutaryl-coenzyme A reductase] [GN:AF1736] [OR:Archaeoglobus fulgidus] [DB:genpept-bct2] [DE:Archaeoglobus fulgidus section 124 of 172 of the complete genome.] [NT:similar to SP:P13702 GB:M29727 GB:M31807 PID:151259] [LE:7093]
[RE:8403] [DI:complement]

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ORF Name	NTID (AAID	NT AA Length Lengt	h score	probability
AI7503000985_2735807_c3_1983	1102	4874	138 45	Ī	
Description					
NO-HIT					
ORF Name	NTID	AAID	NT AA LengthLengt	h score	probability
A17503000985_2739050_c2_1816	1103	4875	927 308	583 1	.2e-56
Description					
sp:[LN:YWBI_BACSU] [AC:P39592] [GN:Y [DE:HYPOTHETICAL TRANSCRIPTIONAL REG [DB:swissprot] >pir:[LN:S39679] [AC:YWBI:protein ipa-24d] [GN:YWBI] [CL [OR:Bacillus subtilis] [DB:pir2] >gp [OR:Bacillus subtilis] [DB:genpept-be [SP:P39592] [LE:24460] [RE:25365] [D [AC:Z99123:AL009126] [GN:YWBI] [FN:UB:DE:Bacillus subtilis complete genome [NT:alternate gene name: ipa-24d; simplice [DI:complement]	ULATOR 1 S39679:0 :probabl :[GI:g41 ct1] [DI I:direct nknown] e (secti	IN THING 370051 Le trance 13948] E:B.sulce IS >gp [OR:Batton 20	K-EPR INTERG PN:transumscription ration [LN:BSGENR] otilis genomatical genom	SENIC REG scription regulator [AC:X73 nic regic 80:g26363 rilis] [Dom 379840	[SION] [SP:P39592] [Tregulator homolog [Signature] [SION] [SP:P39592] [SION] [S
ORF Name	NTID	AAID	<u>NT AA</u> LengthLengt		probability
AI7503000985_2739561_f1_99 Description	1104	4876	1440 479	1505 2	.5e-154
sp:[LN:GLPT_BACSU] [AC:P37948] [GN:G [SP:P37948] [DB:swissprot] >pir:[LN: [PN:glycerol-3-phosphate transport p: [GN:glpT] [CL:hexose phosphate trans [DB:pir2] >gp:[GI:d1034077:g3599636] subtilis] [SR:Bacillus subtilis (strastilis genomic DNA, 70 kb region be [DI:complement] >gp:[GI:g403372] [LN permease] [GN:glpT] [FN:uptake of gl: [DB:genpept-bct1] [DE:B.subtilis glp' permeaseand glycerophosphoryl diester [RE:1649] [DI:direct] >gp:[GI:e118216] [PN:glycerol-3-phosphate permease] [OR:Bacillus subtilis] [DB:genpept-bc 2 of 21): from 194651 to415810.] [NT [RE:40667] [DI:complement]	I40417] rotein g sport pr [LN:ABC ain:168) etween I :BSGLPTC ycerol 3 I and g] r phosph 66:g2632 GN:glpT] ct1] [DE	[AC:I4] [JDT:g] [JDT:g	40417:F69634 lycerol-3-ph uhpT] [OR:E [AC:AB0064 [DB:genpept 23degree.] :Z26522] [PN phate] [OR:E nes for glycerase.] [SE [LN:BSUB0002 uptake of gl	e:S37250 dosphate dacillus (24] [GN: (-bct1) [[LE:3698 d:glycero dacillus derol 3-p (:P37948] d: [AC:Z9 ycerol-3	permease glpT] subtilis] ybeE] [OR:Bacillus DE:Bacillus 4] [RE:38318] 1 3-phosphate subtilis] hosphate [LE:315] 9105:AL009126] -phosphate] te genome (section
ORF Name	NTID	AAID	<u>NT AA</u> LengthLengt	score	probability
AI7503000985_2760930_f1_121	1105	4877		<u>=</u> 	
Description				_	
NO-HIT	<u>0_0</u>				
ORF Name AI7503000985_2767577_c3_2139	NTID 1106	AAID 4878	NT AA LengthLengt	h score	probability
Description			· ·		

ORF Name	NTID	AAID	NI LengthLe	ength score	probability
AI7503000985_2790936_£2_823	1107	4879			3.1e-83
Description sp:[LN:MANA_STRMU] [AC:Q59935] [GN:P [DE:(PMI) (PHOSPHOHEXOMUTASE)] [SP:Q [LN:STRPMI] [AC:D16594] [PN:Mannosep mutans] [SR:Streptococcus mutans (st [DE:S.mutans pmi gene for mannosepho	59935] hosphate rain:GS sphate	[DB:sw e Isom -5) DN isomer	vissprot] merase] [0 MA] [DB:ge mase (comp	>gp:[GI:d GN:pmi] [O enpept-bct plete cds)	1004537:g451216] R:Streptococcus 1] [EC:5.3.1.8]
fructokinase (partial cds).] [LE:241] [RE:1:	191] [DI:direct	=]	
ORF Name	NTID	AAID	<u>NT</u> LengthLe	AA ength	probability
AI7503000985_2814000_c3_2159	1108	4880	156 5		
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> LengthLe	AA ength	probability
A17503000985_2819803_c3_1966	1109	4881	363 12	20 206	1.1e-16
Description					· · · · · · · · · · · · · · · · · · ·
<pre>gp:[GI:e1312399:g3341642] [LN:VCH231 [DB:genpept-bct1] [DE:Vibrio cholera</pre>					
ORF Name	NTID	AAID	NT LengthLe	AA ngth score	probability
A17503000985_2823562_c3_2100 Description	1110	4882	1365 4	909	3.5e-91
pir: [LN:A71026] [AC:A71026] [PN:prohorikoshii] [DB:pir2] >gp:[GI:d10315 [AC:AP000006:AB005215:AB009510:AB009 hypothetical aminotransferase] [GN:Phorikoshii (strain:OT3) DNA, clone:P[DE:Pyrococcus horikoshii OT3 genomi [NT:similar to owl:D5045389 percent [DI:direct]	52:g325 511:AB00 H1501] yrococcu c DNA, 3	7926] 09512: [OR:Py 1s hor 116600	[LN:AP000 AB009513: rococcus ikoshi] 1-1485000	0006] AB009514] horikoshi: DB:genpept	[PN:438aa long i] [SR:Pyrococcus t-bct1] tion(6/7).]
ORF Name	NTID	AAID	<u>NT</u> LengthLe	AA ngth score	probability
A17503000985_2853431_£2_504	1111	4883	195 64		
Description NO-HIT					
ORF Name	NTID	AAID	NT LengthLe	AA ngth score	probability
AI7503000985_2854787_c1_1553	1117	1001	126 41		
	1112	4884		•	

ORF Name	NTID	AAID	NT AA score probability
A17503000985_2867961_c2_1846	1113	4885	963 320 410 2.7e-38
Description			
B] [OR:Bacillus subtilis] [DB:pir2] [AC:AB000617] [PN:YcdH] [GN:ycdH] [O (strain:168 trpC2) DNA] [DB:genpept- degree region, completecds.] [NT:hom [RE:22380] [DI:direct] >gp:[GI:e1182	>gp:[GI R:Bacili bct1] [1 ologue (237:g26] subtili rom 194(:d1023 lus su DE:Bac of adh 32571] s] [DB	btilis] [SR:Bacillus subtilis illus subtilis genomic DNA, 22 to 25 esion protein precursor of] [LE:21421] [LN:BSUB0002] [AC:Z99105:AL009126] :genpept-bct1] [DE:Bacillus subtilis
ORF Name	NTID	AAID	NT AA score probability
A17503000985_29304552_c3_2150	1114	4886	TI38 45
Description			
NO-HIT			
ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000985_29352312_c3_2063	1115	4887	765 254 617 3.1e-60
ABC TRANSPORTER ATP-BINDING PROTEIN >pir:[LN:A70074] [AC:A70074] [PN:AB [GN:yxdL] [CL:ATP-binding cassette	IN IDH : C trans; homolog; O] [AC:I subtilis lis 15] sion pro O] [LN:I [DB:gen; 281to 42; LE:69902 2] [AC:I A1 (Mark e betwee	3'REGIO porter y] [OR D14399] s (stra kb chro betein [BSUB00] pept-be 214814 2] [RE D45912] burg 10 en the	(ATP-binding protein) homolog yxdL] :Bacillus subtilis] [DB:pir2]] [PN:hypothetical protein] [GN:B65F] ain:BGSC 1A1 (168 trpC2)) DNA] comosome segment contains the iol FtsE of E.] [LE:13014] [RE:13787] 21] [AC:Z99124:AL009126] [GN:yxdL] ctl] [DE:Bacillus subtilis complete .] [NT:similar to ABC transporter :70675] [DI:complement]] [GN:yxdL] [OR:Bacillus subtilis] 68; trpC2)) DNA] [DB:genpept-bct1] iol and hut operon,partial and
ORF Name	NTID	AAID	NT AA score probability
A17503000985_29375307_£2_499	1116	4888	 135 44

Description NO-HIT

ORF Name	NTID	AAID	NT AA LengthLength score	probability
A17503000985_29400332_c1_1571	1117	4889	138 309	1.3e-27
Description				
pir:[LN:C69770] [AC:C69770] [PN:hypsubtilis] [DB:pir2] >gp:[GI:d1020028 [OR:Bacillus subtilis] [SR:Bacillus [DE:Bacillus subtilis genome sequence degree.] [NT:FUNCTION UNKNOWN.] [LE:>gp:[GI:e1182404:g2632738] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-b3 of 21): from 402751 to611850.] [LE	:g18812 subtili e, 148 26046] 003] [A	48] [L s (str kb seq [RE:26 C:Z991 E:Baci	N:AB001488] [AC:AB6 ain:168) DNA] [DB:quence of the region 498] [DI:complement .06:AL009126] [GN:yo	O01488] [GN:ydaT] genpept-bct1] nbetween 35 and 47 t] daT] [FN:unknown] lete genome (section
ORF Name	NTID	AAID	NT AA LengthLength score	probability
A17503000985_29493827_£2_872	1118	4890	165 54	
Description				
NO-HIT				
ORF Name	NTID	AAID	NT AA LengthLength score	probability
A17503000985_29503403_f2_788	1119	4891	126 41	
Description				
NO-HIT				
ORF Name	NTID	AAID	NT AA LengthLength score	probability
AI7503000985_29532827_f2_477	1120	4892	243 80	
Description				
NO-HIT				
ORF Name	NTID	AAID	NT AA LengthLength score	probability
A17503000985_29695327_c2_1739	1121	4893	750 249	
Description				
NO-HIT		71.0		
ORF Name	NTID	AAID	NT AA LengthLength	probability
A17503000985_29955003_c2_1894	1122	4894	129 42	
Description				

OKI Name	MIID	MAID	Length	Length	acore	probability
A17503000985_3007827_c2_1943	1123	4895		262		5.8e-59
Description pir: [LN:E69761] [AC:E69761:I40450:S5. [GN:yckK] [CL:lysine-arginine-ornit. [DB:pir2] >gp: [GI:e1182313:g2632647] [FN:unknown] [OR:Bacillus subtilis] genome (section 2 of 21): from 19465 transporter] [LE:215587] [RE:216393] [LN:BSUB0003] [AC:Z99106:AL009126] [GB:genpept-bct1] [DE:Bacillus subtilited [DB:genpept-bct1]] [DE:Bacillus subtilited [DI:complement]] >gp: [GI:d1009629:g18] [glutamine-binding periplasmic] [GN:ycomplement]]	hine-bir [LN:BSU [DB:geng 1 to4158 [DI:com GN:yckK] lis comg ABC tran 05432] [ckK] [OF	ding properties of the propert	protein [AC:2 ct1] [I [NT:sin nt] >gr nknown genome er] [LE 0453] [lus su illus s	n] [OR: 299105: DE:Baci milar t D:[GI:e n] [OR: (secti E:7487] [AC:D50 ubtilis	Bacill AL0091 llus s o glut 118232 Bacill on 3 c [RE:8 453] [] [SR: s DNA	us subtilis] 26] [GN:yckK] ubtilis complete amine ABC 8:g2632662] us subtilis] ff 21): from 402751 293] PN:homologue of Bacillus subtilis for 25-36 degree
ORF Name A17503000985_30078378_c1_1437 Description	NTID 1124			AA Length 54	score	probability
ORF Name A17503000985_3009382_c3_1965 Description NO-HIT	NTID 1125	<u>AAID</u> 4897		AA Length	score	probability
ORF Name A17503000985_30095011_f1_179 Description NO-HIT	NTID	<u>AAID</u> 4898		AA Length	score	probability
ORF Name A17503000985_30133562_c1_1399 Description Sp:[LN:YYBF_BACSU] [AC:P37498] [GN:YN KD PROTEIN IN COTF-TETB INTERGENIC RE [AC:S65991:A70087] [PN:membrane protesistance protein yybF] [OR:Bacillus [LN:BAC180K] [AC:D26185] [PN:unknown] (sub_species:Marburg, strain:168) DNA kilobase region of replication origin pgp:[GI:e1184792:g2636613] [LN:BSUB00] [OR:Bacillus subtilis] [DB:genpept-box 21 of 21): from 3999281to 4214814.] [SP:P37498] [LE:179091] [RE:180305]	(BF] [OR EGION] [cein yyb s subtil [OR:Ba A] [DB:g h.] [LE: 021] [AC ct1] [DE	:BACII SP:P37 F] [GN is] [I cillus enpept 27159] :Z9912 :Bacil lar to	LUS SU 498] [1:yybF B:pir2 subti -bct1] [RE:2 4:AL00 lus su	BTILIS DB:swis] [CL:]] >gp: lis] [S [DE:B 8373] 9126] btilis	DE: ssprot probab [GI:d1 SR:Bac subt [DI:co: [GN:yy compl	<pre>] >pir:[LN:S65991] le antibiotic 005739:g467351] illus subtilis ilis DNA, 180 mplement] bF] [FN:unknown] ete genome (section</pre>
ORF Name A17503000985_302_f2_645 Description		<u>AAID</u> 4900	NT Length	<u>AA</u> Length ⁵	score	probability

A17503000985_30360925_f2_901
Description
sp:[LN:Y357_HAEIN] [AC:P44658] [GN:HI0357] [OR:HAEMOPHILUS INFLUENZAE] [DE:PUTATIVE THIAMINE BIOSYNTHESIS PROTEIN HI0357] [SP:P44658] [DB:swissprot] >pir:[LN:C64063] [AC:C64063] [PN:hypothetical protein HI0357] [OR:Haemophilus influenzae] [DB:pir2] >gp:[GI:g1573325] [LN:U32720] [AC:U32720:L42023] [PN:thiamine biosynthesis protein, putative] [GN:HI0357] [OR:Haemophilus influenzae Rd] [DB:genpept-bct2] [DE:Haemophilus influenzae Rd section 35 of 163 of the complete genome.] [NT:similar to SP:P42883 SP:P43534 SP:P47183] [LE:3006] [RE:3950] [DI:direct]
ORF Name NTID AAID NT AA probability
AI7503000985_30367767_£1_56
Description
pir: [LN:D71235] [AC:D71235] [PN:hypothetical protein PH0142] [GN:PH0142] [OR:Pyrococcus horikoshii] [DB:pir2] >gp:[GI:d1030154:g3256528] [LN:AP000001] [AC:AP000001:AB009465:AB009464:AB009466:AB009467:AB009468:AB009469] [PN:289aa long hypothetical protein] [GN:PH0142] [OR:Pyrococcus horikoshii] [SR:Pyrococcus horikoshii (strain:OT3) DNA] [DB:genpept-bct1] [DE:Pyrococcus horikoshii OT3 genomic DNA, 1-28700 nt. position (1/7).] [NT:motif=soybean trypsin inhibitor (Kunitz) protease] [LE:124718 [RE:125587] [DI:direct]
AI7503000985_30470325_c3_2181
pir:[LN:A70027] [AC:A70027] [PN:conserved hypothetical protein yvaC] [GN:yvaC] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1186043:g2635868] [LN:BSUB0018] [AC:Z99121:AL009126] [GN:yvaC] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 18 of 21): from 3399551to 3609060.] [NT:similar to hypothetical proteins] [LE:45749] [RE:47644] [DI:complement]
ORF Name NTID AAID NT AA score probability
A17503000985_30651577_c3_2155 1132 4904 612 203 88 0.00087
Description
pir:[LN:G70065] [AC:G70065] [PN:hypothetical protein ywpE] [GN:ywpE] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1184540:g2636159] [LN:BSUB0019] [AC:Z99122:AL009126] [GN:ywpE] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 19 of 21): from 3597091to 3809700.] [LE:143698] [RE:144006] [DI:complement] >gp:[GI:e289144:g1763706] [LN:BSZ83337] [AC:Z83337] [GN:ywpE] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:B.subtilis mbl, flh[O,P], rapD, ywp[B,C,D,E,F,G,H,I,J] and ywqAgenes.] [LE:5315] [RE:5623] [DI:direct] >gp:[GI:e1184540:g2636159] [LN:BSUB0019] [AC:Z99122:AL009126] [GN:ywpE] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept] [DE:Bacillus subtilis complete genome (section 19 of 21): from 3597091to 3809700.] [LE:143698] [RE:144006] [DI:complement]
ORF Name NTID AAID NT AA score probability A17503000985_31490687_f3_1096
Description NO-HIT

ORF Name

AAID NT AA score probability

ORF Name	NTID	AAID NT AA score probability
AI7503000985_31517587_c2_1663	1134	4906 129 42
Description		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000985_3157062_c1_1475	1135	4907 174 57
Description		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000985_31693_c2_1716	1136	4908 699 232 128 7.1e-08
Description		
pir:[LN:JH0364] [AC:JH0364] [PN:hyp [OR:Streptococcus pyogenes] [DB:pir2		al protein 176 (SAGP 5' region)]
ORF Name	NTID	AAID NT AA score probability
A17503000985_31720942_f2_813	1137	4909 576 191
Description		
NO-HIT	and an exercise from	
ORF Name	NTID	AAID <u>NT AA</u> score probability
AI7503000985_31755012_c1_1340	1138	4910 978 325 886 9.6e-89
Description		
[CL:glucose transport protein] [OR:B >gp:[GI:e1184489:g2636109] [LN:BSUB0	acillus 019] [A	C:Z99122:AL009126] [GN:ywtG] [FN:unknown]
	[NT:sim	E:Bacillus subtilis complete genome (section ilar to metabolite transport protein] GI:e308095:q1894771] [LN:BSZ92954]
[AC:Z92954] [GN:ywtG] [OR:Bacillus s	ubtilis] [DB:genpept-bct1] [DE:B.subtilis
yws[A,B,C,D,E,F,G] and gerBC genes.]		oduct highly similar to metabolite >gp:[GI:e1184489:g2636109] [LN:BSUB0019]
[AC:Z99122:AL009126] [GN:ywtG] [FN:u	nknown]	[OR:Bacillus subtilis] [DB:genpept]
		ion 19 of 21): from 3597091to 3809700.]
[NT:Similar to metabolite transport	protein] [LE:94500] [RE:95873] [DI:complement]
ORF Name	NTID	AAID <u>NT AA</u> score probability
A17503000985_3182927_c2_1673	1139	4911 2082 693 3598 0.0
Description [77]		
<pre>gp:[GI:g2981225] [LN:AF053006] [AC:A [OR:Staphylococcus epidermidis] [DB: precursor (geh1) gene, completecds.]</pre>	genpept	-bct2] [DE:Staphylococcus epidermidis lipase
ORF Name	NTID	AAID NT AA score probability
A17503000985_32205143_c2_1642	1140	4912 168 55
Description		
NO-HIT		

ORF Name	NTID	AAID NT AA score probability
A17503000985_32220202_f2_708	1141	4913 123 40
Description	<u> </u>	
NO-HIT		
		NT AA
ORF Name	NTID	AAID LengthLength score probability
A17503000985_32221012_c1_1567	1142	4914 471 156 114 4.6e-06
Description		
	MSPORTE U90545] pept-pr	(R 4)] [SP:000476] [DB:swissprot] [PN:sodium phosphate transporter] [GN:NPT4] (i2) [DE:Human sodium phosphate transporter
ORF Name	NTID	AAID NT AA score probability
AI7503000985_32614078_f2_539	1143	4915 150 49
Description		
NO-HIT		
		NTP 7.7
ORF Name	NTID	AAID LengthLength score probability
AI7503000985_32664093_£2_568	1144	4916 627 208 142 6.7e-10
Description	L	
[CL:probable thiamin-phosphate pyrophomology] [OR:Aquifex aeolicus] [DB: [AC:AE000736:AE000657] [PN:thiamine	hosphor pir2] > phospha fex aeo	te synthase] [GN:thiE1] [OR:Aquifex licus section 68 of 109 of the complete
ORF Name	NTID	AAID NT AA probability LengthLength score probability
AI7503000985_33203385_c3_1997	1145	4917 597 198 967 2.5e-97
Description		
epidermidis] [DB:genpept-bct2] [DE:S	taphylo	[PN:IcaR] [GN:icaR] [OR:Staphylococcus coccus epidermidis operon mediating aB, and IcaC genes, complete cds.] [LE:39]
ORF Name	NTID	AAID NT AA score probability
A17503000985_33209682_c2_1718	1146	4918 123 40
Description		
NO-HTT		

ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	score	probability
A17503000985_33211092_£3_1123	1147	4919	471	156	72	0.030
Description sp:[LN:VL02_VARV] [AC:P33041] [GN:L2E [SP:P33041] [DB:swissprot] >pir:[LN:Sprotein] [CL:vaccinia virus F3 protein [LN:S55844] [AC:S55844] [GN:L2R] [OR: India-1967] [DB:genpept-vrl] [DE:G9R. 19 genes,18029 nt].] [NT:This sequence [DI:direct] >gp:[GI:g438992] [LN:VARC [SR:Variola major virus (strain Banglamajor virus	333088] in] [OR VariolaH7R ce comes CG] [AC ladesh-: complet 519] [DB 58886] virus] it) gene [LN:VVI Garcia-:	[AC:Sivarion in the content of the c	33088:H la viru r virus la majo Fig. 2 9] [OR: DNA] [D DME.] [ct] >gp ot-vrl] rect] > enpept- SP:P330 AC:X762	36844 s] [DE] [SR: r viru .] [LE Variol B:genp NT:hom :[GI:g [DE:V gp:[GI vrl] [41] [I 67] [C L,N4R	PN P	LL2R protein:M2R >gp:[GI:g262433] la major virus dia-1967, Genomic, [RE:2123] or virus] fl] [DE:Variola of vaccinia virus l] [LN:VVCGAA] a virus DNA complete 50] [LN:VVHINDQKH] riola virus 08] [RE:18771] tola virus]
ORF Name AI7503000985_33219007_t3_1152 Description NO-HIT	NTID 1148	AATD	<u>NT</u> Length	<u>AA</u>	score	probability
ORF Name AI7503000985_33241093_c2_1806 Description pir:[LN:A69759] [AC:A69759] [PN:1-py [GN:ycgN] [CL:aldehyde dehydrogenase [OR:Bacillus subtilis] [DB:pir2] >gp: [AC:Z99105:AL009126] [GN:ycgN] [FN:un [DE:Bacillus subtilis complete genome [NT:similar to 1-pyrroline-5-carboxyl [DI:direct] >gp:[GI:d1009590:g1805393 to] [GN:ycgN] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtil theamyE-srfA region, complete cds.] [e (NAD+) [GI:el] known] e (secti ate del [SR:Bac is DNA	4921 e-5-can :aldel 182273: [OR:Baion 2 conydroge 050453] cillus for 25	rboxyla nyde del g26326 acillus of 21): enase] [AC:D subtil	te deh hydrog 07] [L subti from [LE:15 50453] is (st gree r	ydroge enase N:BSUE lis] (194651 0401] [PN:6 rain:1	homology] 30002] [DB:genpept-bct1] 10415810.] [RE:151948] 8% identity protein 168 trpC2) DNA] containing
ORF Name AI7503000985_33242842_c1_1579 Description pir:[LN:S59797] [AC:S59797] [PN:hypo D9798.1] [CL:unassigned DEAD/H box he [OR:Saccharomyces cerevisiae] [DB:pir	licases	4922 al prot	ein YD	956 R332w:	hypoth	-
	NTID	<u>AAID</u> 4923	NT LengthI	<u>AA</u> Length 51	score	probability

ORF Name	NTID	AAID NT AA score probability
A17503000985_33304082_f1_387	1152	4924 129 42
Description	لـــــا	
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000985_33359381_c2_1697	1153	4925 210 69
Description		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000985_33391337_c2_1891	1154	4926 864 287 168 2.1e-12
Description		
		PN:pXO1-85] [OR:Bacillus anthracis]
[DB:genpept-bct2] [DE:Bacilius anthr [LE:99636] [RE:100319] [DI:direct]	acıs vı:	rulence plasmid PX01, complete sequence.]
		NT AA
ORF Name	NTID	AAID LengthLength score probability
A17503000985_33595087_c2_1645	1155	4927 141 46
Description NO-HIT		
NO-III		
ORF Name	NTID	AAID <u>NT AA</u> score probability LengthLength
A17503000985_33620176_c3_2091	1156	4928 858 285 494 3.3e-47
Description		
gp:[GI:d1020251:g1943993] [LN:AB0018 aureus] [SR:Staphylococcus aureus (s		:AB001896] [GN:orf30] [OR:Staphylococcus
_ -		peron, complete cds.] [LE:1501] [RE:2310]
[DI:direct]		
ORF Name	NTID	AAID NT AA score probability
A17503000985_33837817_c1_1431	1157	4929 129 42
Description		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
AI7503000985_33985007_f3_1266	1158	4930 153 50
Description		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
AI7503000985 33988778 ±2 888	1159	LengthLength
Description		
pir:[LN:JC4511] [AC:JC4511] [PN:pyr		yl-peptidase I,:bacterial pyrrolidone
carboxyl peptidase (PYRase)] [GN:pcp		
[DB:pir2] >gp:[GI:g790573] [LN:SAU19 peptidase] [GN:pcp] [OR:Staphylococc		I:U19770] [PN:pyrrolidone carboxyl us] [DB:genpept-bctl] [DE:Staphylococcus
aureus pyrrolidone carboxyl peptidas		gene, complete cds.] [NT:pyrase] [LE:204]
[RE:842] [DI:direct]		

ORF Name	NTID	AAID	NT AA score probability
A17503000985_34016937_c1_1600	1160	4932	
Description			
	glycine mology] AF009352 llus sul ne protein system] 018] [AG puCA] [1 :genpept	betai [OR:B 2] [PN btilis ein (o (opuC] [LE: C:Z991 FN:hig t-bct1	ine/proline transport protein proV: Bacillus subtilis] [DB:pir2] N:ATPase] [GN:opuCA] [OR:Bacillus S osmoprotectant transport system OpuC DpuCB), osmoprotectantbinding protein CD)genes, complete cds.] [NT:OpuCA; 1860] [RE:2002] [DI:direct] 121:AL009126] [PN:glycine 13 [DE:Bacillus subtilis complete
ORF Name	NTID	AAID	NT AA score probability
A17503000985_34021912_c2_1736	1161	4933	
Description			
NO-HIT			
ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000985_3402312_c1_1529	1162	4934	1761 586 1316 2.6e-134
Description			
	n:thiami :[GI:d10 :Bacillu lis gend :SIMILAM p:[GI:e1 nknown] e (secti	ine py: 020024 us subsome sec R TO P 1182400 [OR:Ba	rophosphate-binding domain homology] 4:g1881244] [LN:AB001488] [AC:AB001488] 5tilis (strain:168) DNA] 6quence, 148 kb sequence of the 7YRUVATE OXIDASE AND ACETOLACTATE] 70:g2632734] [LN:BSUB0003] 73acillus subtilis] [DB:genpept-bct1] 70 of 21): from 402751 to611850.]
ORF Name	NTID	AAID	NT AA probability
A17503000985_34033563_c1_1627	1163	4935	327 108 388 5.7e-36
<pre>product] [LE:2589] [RE:2903] [DI:dire [AC:AF029224:AF029225] [PN:NirD] [GN [DB:genpept-bct2] [DE:Staphylococcus</pre>	nce 1 fr ect] >gr :nirD] carnosu	com Pato: [GI:g [OR:State as nir	tent EP0805205.] [NT:unnamed protein g4433640] [LN:AF029224]
ORF Name	NTID	AAID	NT AA score probability
A17503000985_34094136_f1_408	1164	4936	156 51
Description NO-HIT			

ORF Name	NTID	AAID	NT LengthLe	AA ength s	core	probability
A17503000985_34157807_£2_816	1165	4937	147 48	8		
Description	<u> </u>					
NO-HIT						
ORF Name	NTID	AAID	NT Length Le	AA ength	core	probability
A17503000985_34194002_c2_1941	1166	4938	711 23	36	557 1	.8e-64
Description sp:[LN:PMGY_ZYMMO] [AC:P30798] [GN:P [DE:(BPG-DEPENDENT PGAM)] [SP:P30798 [PN:phosphoglycerate mutase,] [CL:ph homology] [OR:Zymomonas mobilis] [EC [AC:L09651] [PN:phosphoglyceromutase mobilis (strain CP4) DNA] [DB:genpep (pgm) gene, complete cds, and 2-hydro [LE:317] [RE:1003] [DI:direct]	DB:stoophogly :5.4.2.3] [GN:pothersection]	wisspr ycerat 1] [DB gm] [O [DE:Z	ot] >pir: e mutase: :pir2] >g R:Zymomor ymomonas	:[LN:C :phosp gp:[GI nas mo mobil	40649] hoglyc :g1556 bilis] is pho	[AC:C40649] erate mutase 11] [LN:ZMOPGMA] [SR:Zymomonas esphoglyceromutase
ORF Name	NTID	AAID	NT LengthLe	AA ength	core	probability
A17503000985_34197318_£3_1128	1167	4939] 903 30	00 1	117 3	.2e-113
Description	<u></u>					
<pre>pir:[LN:A49943] [AC:A49943:S33358] [OR:Staphylococcus carnosus] [EC:4.1 [AC:X71729] [PN:fructose-bisphosphat [DB:genpept-bct1] [EC:4.1.2.13] [DE: [RE:1557] [DI:direct]</pre>	.2.13] e aldola	[DB:pi ase] [r2] >gp:[GN:fda] [[GI:g2: [OR:St	97874] aphylo	[LN:SCFDA]
ORF Name	NTID	AAID	<u>NT</u> LengthLe	AA ength	core	probability
AI7503000985_34273436_f2_498	1168	4940	855 28	84	372 2	.8e-34
Description		<u></u>	JL L			
sp:[LN:YQJG_BACSU] [AC:P54544] [GN:Y [SP:P54544] [DB:swissprot] >pir:[LN:homolog yqjG] [GN:yqjG] [CL:stage I protein homology] [OR:Bacillus subti [LN:BACJH642] [AC:D84432:D82370] [PN (strain:JH642(trpC2 PheA1)) DNA] [DB region containing skin element.] [LE >gp:[GI:e1185657:g2634823] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-b 13 of 21): from 2395261to 2613730.] [SP:P54544] [LE:87873] [RE:88700] [D	G69963] II sport lis] [DI :YqjG] :genpept :234919] O13] [AC ct1] [DI	[AC:Gulations: AC:Gulations: A	69963] [n protein] >gp:[GI cillus su] [DE:Bac 235746] [16:AL0091 llus subt	[PN:li] n:stage I:d101: ubtilis cillus [DI:con 126] [6 cillis e	poprot e III 3293:g s] [SR subti mpleme GN:yqj comple	sporulation 1303958] :Bacillus subtilis lis DNA, 283 Kb nt] G] [FN:unknown] te genome (section
ORF Name	NTID	AAID	NT LengthLe	AA ngth	core	probability
AI7503000985_34385012_£2_465	1169	4941	1362 45		64 6	.0e-40
Description						
pir:[LN:S62194] [AC:S62194] [PN:hypportein] [OR:Methanosarcina barkeri] [AC:X93084] [GN:orf4] [OR:MethanosarcfmdF, fmdA, fmdC, fmdD, fmdB, orf4, [DI:complement]	DB:piː] cina baː	:2] >g keri]	p:[GI:e21 [DB:genp	12291:g	g11249 ct1] [:	57] [LN:MBFMDSUBS] DE:M.barkeri fmdE,

ORF Name	NTID	AAID Length Length score probability
A17503000985_34410843_c2_1690	1170	4942 1431 476
Description		
NO-HIT		
		NT AA
ORF Name	NTID	AAID LengthLength score probability
AI7503000985_34412952_c3_2210	1171	4943 783 260 580 2.6e-56
Description		
sp:[LN:YBBM_ECOLI] [AC:P77307] [GN:Y PROTEIN IN USHA-TESA INTERGENIC REGI		R:ESCHERICHIA COLI] [DE:HYPOTHETICAL 28.2 KD:P77307] [DB:swissprot]
ORF Name	NTID	AAID NT AA score probability
A17503000985_34414192_c3_2229	1172	4944 831 276 113 0.00012
Description		
homolog MTH178:Icc related protein] phosphodiesterase cpdA: 3',5'-cyclic homology:phosphoesterase core homolo [EC:3.1.4.17] [DB:pir1] >gp:[GI:g262 related protein] [GN:MTH178] [OR:Met [DB:genpept-bct1] [DE:Methanobacteri	[GN:MTH -nucleo gy] [OR 1221] [i hanobac um ther plete g	tide phosphodiesterase cpdA :Methanobacterium thermoautotrophicum] LN:AE000805] [AC:AE000805:AE000666] [PN:Icc terium thermoautotrophicum] moautotrophicum from bases 114371 to enome.] [NT:Function Code:10.02 - Metabolism
ORF Name	NTID	AAID NT AA score probability
A17503000985_34430428_c2_1949 Description	1173	4945 375 124 83 0.0051
subtilis] [DB:pir2] >gp:[GI:d1020089 [OR:Bacillus subtilis] [SR:Bacillus [DE:Bacillus subtilis genome sequenc degree.] [NT:FUNCTION UNKNOWN.] [LE: >gp:[GI:e1182465:g2632799] [LN:BSUB0	:g188130 subtili: e, 148 1 78831] 003] [Ad ct1] [Di	C:Z99106:AL009126] [GN:yddJ] [FN:unknown] E:Bacillus subtilis complete genome (section
ORF Name	NTID	AAID NT AA score probability
AI7503000985_34554692_f1_62	1174	4946 474 157 403 1.5e-37
Description		
<pre>gp:[GI:g1575025] [LN:SAU52961] [AC:U [OR:Staphylococcus aureus] [SR:Staph [DB:genpept-bct2] [DE:Staphylococcus</pre>	ylococci aureus	
ORF Name	NTID	AAID NT AA score probability
AI7503000985 34585317 c2 1938	1175	LengthLength 130 2.6e-05
Description		
gp:[GI:g454844] [LN:SCMP48EGG] [AC:M		[OR:Schistosoma mansoni] [SR:Schistosoma
		DB:genpept-inv1] [DE:Schistosoma mansoni [DE:ORF 3] [LE:687] [RE:1868] [DI:complement]
bro caanings brocciii aciic, combiere	LI	AI.OKI DI [HE.OO/] [KE.IOOO] [DI:COMPIEMENE]

ORF Name	NTID AAID NT AA score probability
A17503000985_34617937_£3_1227	1176 4948 525 174 223 1.3e-17
Description	
[OR:Dictyostelium discoideum] [DB:ge	AF051898] [PN:coronin binding protein] [GN:DB10] enpept-inv1] [DE:Dictyostelium discoideum coronin ecds.] [LE:108] [RE:1790] [DI:direct]
ORF Name	NTID AAID NT AA score probability
AI7503000985_34647150_c1_1439	1177 4949 1356 451 274 1.6e-23
Description	
[GN:shp3] [OR:Litomosoides sigmodont microfilarial sheath proteins SHP3a([PN:microfilarial sheath protein SHP3] [Is] [DB:genpept-invl] [DE:Litomosoides sigmodontis shp3a) and SHP3 (shp3) genes, complete cds.] [Shp3 genes from] [LE:7991:8260] [RE:8047:9219]
ORF Name	NTID AAID NT AA probability LengthLength
A17503000985_35193950_c2_1811	1178 4950 489 162
Description	
NO-HIT	
ORF Name	NTID AAID NT AA score probability
AI7503000985_35317188_c2_1684	1179 4951 6414 2137 2779 2.4e-289
[GN:fap1] [OR:Streptococcus parasang	F100426] [PN:fimbriae-associated protein Fap1] ruinis] [DB:genpept-bct2] [DE:Streptococcus ein Fap1 (fap1)gene, complete cds.] [NT:invovled in ted] [LE:284] [RE:7996] [DI:direct]
ORF Name	NTID AAID NT AA score probability
A17503000985_35433438_f3_1030	1180 4952 1137 378 420 2.3e-39
Description	
[DE:GLYCINE OXIDASE,] [SP:O31616] [DI:PN:sarcosine oxidase homolog yjbR] >gp:[GI:e1183187:g2633521] [LN:BSUB06] [OR:Bacillus subtilis] [DB:genpept-b6]	UBR] [OR:BACILLUS SUBTILIS] [EC:1.5.3] (B:swissprot] >pir:[LN:B69845] [AC:B69845] [GN:yjbR] [OR:Bacillus subtilis] [DB:pir2] 007] [AC:Z99110:AL009126] [GN:yjbR] [FN:unknown] (ct1] [DE:Bacillus subtilis complete genome (section NT:similar to sarcosine oxidase] [SP:O31616]

NT AAORF Name NTID AAID score probability LengthLength A17503000985 35449093_c2 1668 1181 4953 831 276 103 0.0069 Description sp:[LN:YHI3 LACLA] [AC:Q02147] [OR:LACTOCOCCUS LACTIS] [SR:,SUBSPLACTIS:STREPTOCOCCUS LACTIS] [DE:HYPOTHETICAL 38.0 KD PROTEIN IN HISC-HISG INTERGENIC REGION (ORF3)] [SP:Q02147] [DB:swissprot] >pir:[LN:C45734] [AC:C45734] [PN:histidyl-tRNA synthetase homolog] [OR:Lactococcus lactis subsp. lactis] [DB:pir2] >gp:[GI:g2565140] [LN:LLU92974] [AC:U92974:M90760:M90761] [PN:unknown] [OR:Lactococcus lactis] [DB:genpept-bct1] [DE:Lactococcus lactis unknown gene, partial cds, and HisC (hisC), unknown, HisG (hisG), unknown, HisB (hisB), unknown, HisH (hish), HisA (hisA), HisF (hisF), HisIE (hisIE), unknown, unknown, LeuA(leuA), LeuB (leuB), LeuC (leuC), LeuD (leuD), unknown, IlvD(ilvD), IlvB (ilvB), IlvN, IlvC (ilvC), IlvA (ilvA), AldB (aldB) and aldR (aldR) genes, complete cds.] [NT:ORF2; potential regulator; similar to histidyl-tRNA] [LE:2146] [RE:3132] [DI:direct] NT AΑ Length Length score ORF Name NTID AAID probability AI7503000985 35687 f2 812 4954 162 1182 153 Description NO-HIT NTAAORF Name NTID AAID <u>score</u> probability LengthLength AI7503000985 35817137 f3 1254 1183 4955 156 51 Description NO-HIT NTAΑ ORF Name NTID AAID score probability LengthLength A17503000985_35993802_c2_1685 1184 4956 1584 527 Description NO-HIT NT AAAAID ORF Name NTID score probability LengthLength A17503000985_35995316_£1_393 1185 4957 1257 418 2160 9.6e-224 Description

pir:[LN:S77608] [AC:S77608] [PN:probable intercellular adhesion protein A:glycosyltransferase:icaA protein] [GN:icaA] [OR:Staphylococcus epidermidis] [DB:pir2] >gp:[GI:g1161380] [LN:SEU43366] [AC:U43366] [PN:IcaA] [GN:icaA] [OR:Staphylococcus epidermidis] [DB:genpept-bct2] [DE:Staphylococcus epidermidis operon mediating intercellular adhesion: IcaR, IcaA, IcaB, IcaB, and IcaC genes, complete cds.] [LE:761] [RE:1999] [DI:direct]

NT ORF Name NTID AAID probability

AI7503000985 36127302_c2_1875

LengthLength

4958 888

295

1058

Description

sp:[LN:GTAB BACSU] [AC:Q05852] [GN:GTAB] [OR:BACILLUS SUBTILIS] [EC:2.7.7.9] [DE:(GENERAL STRESS PROTEIN 33) (GSP33)] [SP:Q05852] [DB:swissprot] >pir:[LN:A40650] [AC:A40650:B69638] [PN:UTP--glucose-1-phosphate uridylyltransferase,:UDP-glucose pyrophosphorylase] [GN:gtaB] [CL:Escherichia coli UTP--glucose-1-phosphate uridylyltransferase] [OR:Bacillus subtilis] [EC:2.7.7.9] [DB:pir1] >gp:[GI:g289287] [LN:BACGTABX] [AC:L12272] [PN:UDP-glucose pyrophosphorylase] [GN:gtaB] [OR:Bacillus subtilis] [SR:Bacillus subtilis (strain 168, sub species Marburg) DNA] [DB:qenpept-bct1] [EC:2.7.7.9] [DE:Bacillus subtilis UDP-qlucose pyrophosphorylase (gtaB) gene, complete cds.] [NT:similar to UDP-glucose pyrophosphorylase of] [LE:120] [RE:998] [DI:direct] >gp:[GI:g405623] [LN:BSLYTGTA] [AC:Z22516] [PN:UDP-glucose pyrophosphorylase] [GN:gtaB] [OR:Bacillus subtilis] [DB:genpept-bct1] [EC:2.7.7.9] [DE:B.subtilis lytR, orfX, and gtaB genes.] [NT:similar to other procaryotic UDP-glucose] [SP:Q05852] [LE:1581] [RE:2459] [DI:direct] >qp:[GI:e1184473:q2636093] [LN:BSUB0019] [AC:Z99122:AL009126] [PN:UTP-glucose-1-phosphate uridylyltransferase] [GN:gtaB] [FN:glucosylation of teichoic acid] [OR:Bacillus subtilis] [DB:genpept-bct1] [EC:2.7.7.9] [DE:Bacillus subtilis complete genome (section 19 of 21): from 3597091to 3809700.] [SP:Q05852] [LE:67599] [RE:68477] [DI:direct] >gp:[GI:e1184473:g2636093] [LN:BSUB0019] [AC:Z99122:AL009126] [PN:UTP-glucose-1-phosphate uridylyltransferase] [GN:gtaB] [FN:glucosylation of teichoic acid] [OR:Bacillus subtilis] [DB:genpept] [EC:2.7.7.9] [DE:Bacillus subtilis complete genome (section 19 of 21): from 3597091to 3809700.] [SP:Q05852] [LE:67599] [RE:68477] [DI:direct]

ORF Name	NTID	AAID	NT Length1	<u>AA</u> Length	score	probability
A17503000985_36134678_c3_2220	1187	4959	699	232	674	2.8e-66

Description

pir:[LN:F69670] [AC:F69670] [PN:glycine betaine/carnitine/choline ABC transporter (membrane p) opuCD] [GN:opuCD] [CL:glycine betaine/carnitine/choline ABC transporter] [OR:Bacillus subtilis] [DB:pir2] >qp:[GI:q2271392] [LN:AF009352] [AC:AF009352] [PN:transmembrane protein] [GN:opuCD] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis osmoprotectant transport system OpuC includingATPase (opuCA), transmembrane protein (opuCB), osmoprotectantbinding protein precursor (opuCC) and transmembrane protein (opuCD)genes, complete cds.] [NT:OpuCD; part of the osmoprotectant transport system] [LE:3627] [RE:4316] [DI:direct] >gp:[GI:e1186068:g2635893] [LN:BSUB0018] [AC:Z99121:AL009126] [PN:qlycine betaine/carnitine/choline ABC] [GN:opuCD] [FN:high affinity transport of glycine betaine,] [OR:Bacillus subtilis] [DB:qenpept-bct1] [DE:Bacillus subtilis complete genome (section 18 of 21): from 3399551to 3609060.] [NT:alternate gene name: yvbB] [LE:67059] [RE:67748] [DI:complement]

NT AΑ ORF Name NTID AAID score probability LengthLength A17503000985_36142510_c1_1394 1188 132 4960 Description

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probabi	lity	
AI7503000985_36151692_c2_1840	1189	4961	681	226		5.8e-66]
Description		L	لــــال		اــــــا			1
sp:[LN:YFKO_BACSU] [AC:O34475] [GN:Y [DE:PUTATIVE NAD(P)H NITROREDUCTASE [AC:B69809] [PN:NAD(P)H-flavin oxid [CL:nitroreductase] [OR:Bacillus sub [LN:BSUB0005] [AC:Z99108:AL009126] [[DB:genpept-bct1] [DE:Bacillus subti to1011250.] [NT:similar to NAD(P)H-f [RE:51764] [DI:direct] >gp:[GI:d1024 [OR:Bacillus subtilis] [SR:Bacillus [DE:Bacillus subtilis genomic DNA, 7 [DI:complement]	YFKO,] doreduct tilis] [GN:yfkO lis com lavin o 284:g26 subtili	[SP:03-ase hor [DB:pi:] [FN:1] plete (xidored 26827] s (stra	4475] [nolog y r2] >gp nknown genome ductase [LN:D8	[DB:swi rfkO] [p:[GI:e i] [OR: (secti e] [SP: 3967]	Ssprot GN:yfk 118277 Bacill on 5 c O34475 [AC:D8	:] >pir:[L :0] /3:g263310 .us subtil of 21): fr [LE:510 33967] [PN 3:genpept-	7] is] om 80282 99] :YfkO]	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probabi	lity	
A17503000985_36205285_c2_1717	1190	4962	780	259	284	6.0e-25		i
Description sp:[LN:XYNC_CALSA] [AC:P23553] [GN:X [SR:,CALDICELLULOSIRUPTOR SACCHAROLY (ACETYLXYLOSIDASE)] [SP:P23553] [DB: [PN:acetylesterase, (XynC)] [OR:Cald >gp:[GI:g144297] [LN:CDCXYNAB] [AC:M [SR:C.saccharolyticum DNA, clone pNZ xylanase A (XynA), beta-xylosidase (cds.] [NT:acetyl esterase (XynC)] [L [LN:AF005383] [AC:AF005383] [PN:acet saccharolyticus] [DB:genpept-bct2] [transport protein(XynG), putative tr (XynE), xylanase (XynD), xylanase (X genes, complete cds.] [LE:13673] [RE	TICUS] swisspr locellum [34459] [1400] [[XynB) a [E:1257] [Ylxylos [DE:Cald cansport [YnA), a	[EC:3.] ot] >p: saccha [OR:Cal DB:geny ndacety [RE:20 idase] icellul protes cetylxy	ir:[LN: arolyti dicell bept-bc cl este 057] [D [GN:Xy tosirup in (Xyn closida	[DE:AC B37202 cum] [ulosir t1] [D rase (DI:dire mC] [O tor sa H), xy	ETYL E [AC: EC:3.1 uptor E:C.sa XynC) ct] >g R:Cald ccharc lanase	STERASE, B37202] .1.6] [DB saccharol ccharolyt genes, co p:[GI:g26 icellulos lyticus p (XynF),x	yticus] icum mplete 45420] iruptor utative ylanase	
ORF Name	NTID	AAID	<u>NT</u> Lengthi	<u>AA</u> Length	<u>score</u>	probabi	lity	
A17503000985_36214052_c3_1981	1191	4963	999	332	169	1.3e-10		
Description sp:[LN:Y31K_SULAC] [AC:P46218] [OR:S PROTEIN] [SP:P46218] [DB:swissprot] [OR:Sulfolobus acidocaldarius] [DB:g polymerase subunit homolog gene,comp Number P20435:] [LE:61] [RE:888] [DI	>gp:[GI enpept-l	:g45826 bct1] (s.] [N]	5] [LN DE:Sul	:SAU05 folobu	664] [s acid	AC:U05664 ocaldariu] s RNA	
ORF Name	NTID	AAID	<u>NT</u> Lengthl	<u>AA</u> Length	score	probabil	lity	
A17503000985_36225250_c2_1711	1192	4964	1542	513	2622	1.1e-272		
Description								
sp:[LN:SEPA_STAEP] [AC:P43148] [GN:S [DE:EXTRACELLULAR ELASTASE PRECURSOR >pir:[LN:A40659] [AC:A40659] [PN:el [EC:3.4.24] [DB:pir2] >gp:[GI:g396	, (SEPP astase,	l)] [SE SepP1]	P:P4314:	8] [DB taphyl	:swiss ococcu	prot] s epiderm:	idis]]

[GN:SepP1] [OR:Staphylococcus epidermidis] [DB:genpept-bct1] [DE:S.epidermis gene for

protease.] [SP:P43148] [LE:164] [RE:1687] [DI:direct]

ORF Name	NTID	AAID	NT AA LengthLeng	th score	probability
A17503000985_36228252_c3_1990	1193	4965	777 258	678	1.1e-66
Description					
sp:[LN:HIS6_BACSU] [AC:O34727] [GN:H (CYCLASE)] [SP:O34727] [DB:swissprot cyclase-like protein hisF] [GN:hisF [DB:pir2] >gp:[GI:e1186175:g2636000] cyclase-like protein] [GN:hisF] [FN:[OR:Bacillus subtilis] [DB:genpept-b18 of 21): from 3399551to 3609060.] >gp:[GI:g2618870] [LN:AF017113] [AC:subtilis] [DB:genpept-bct2] [DE:Baci[NT:HisF protein] [LE:44003] [RE:447] >pir:] [CL:c [LN:BS synthes ct1] [D [SP:O34 AF01711 llus su	[LN:B6 yclase UB0018 is of E:Baci 727] [727] [Btilis	9641] [AC:E hisF] [OR: left] [AC:Z9912] D-erythro-i llus subtil LE:183072] cyclase] [300-304 de	Bacillus 1:AL0091 midazole is compl [RE:1838 GN:hisF]	[PN:HisF subtilis] 26] [PN:HisF glycerol] ete genome (section 30] [DI:complement] [OR:Bacillus
ORF Name	NTID	AAID	NT AA LengthLeng		probability
AI7503000985_36366326_c2_1646	1194	4966	771 256	409	3.4e-38
Description					
pir: [LN:B69906] [AC:B69906] [PN:ran coli rarD protein] [OR:Bacillus subt [LN:BSUB0011] [AC:Z99114:AL009126] [DB:genpept-bct1] [DE:Bacillus subti 2000171to 2207900.] [NT:similar to h [DI:complement] >gp:[GI:g3169320] [L [FN:unknown] [OR:Bacillus subtilis] (yojA), YojB (yojB), YojC (yojC), Yo YojH (yojH), YojI(yojI), YojJ (yojJ) YojN(yojN), and YojO (yojO) genes, complement [NT:similar to Escherichia coli RarD]	ilis] [] GN:yojE lis com ypothet N:AF026 [DB:gen jD(yojD , YojK omplete	DB:pir] [FN: plete ical p 147] [pept-b), Yoj (yojK) cds;	2] >gp:[GI: unknown] [C genome (sec roteins] [L AC:AF026147 ct2] [DE:Ba E (yojE), Y , YojL (yoj and OdhA (o	e1185420 R:Bacill tion 11 E:122204] [PN:Yo cillus s ojf (yoj L), YojM dhA) gen	<pre>:g2634341] us subtilis] of 21): from] [RE:122995] jD] [GN:yojD] ubtilis YojA F), YojG (yojG), (yojM), e,partial cds.]</pre>
ORF Name	NTID	AAID	NT AA Length Leng		probability
AI7503000985_36617832_c1_1459	1195	4967	801 266	758	3.5e-75
Description					
sp:[LN:SUMT_BACME] [AC:P29928] [GN:COME [DE:METHYLASE) (SUMT) (UROPORPHYRINO [DB:swissprot] >pir:[LN:A42479] [AC:uroporphyrinogen III methyltransferamethyltransferase] [OR:Bacillus megaterium] [SR:Bacillus megaterium] [SR:Bacillus megaterium] S-adenosy-L-methionine:uroporphyrinogeds.] [NT:ATCC #1078] [LE:119] [RE:8]	GEN III A42479 [se] [CL terium] ine:urop DNA] [D] gen III	METHY [PN: S-ade DB:p porphy B:genp methyl	LASE) (UROM S-adenosyl- nosyl-L-met ir2] >gp:[G rinogen III ept-bct1] [transferase)] [SP:P L-methio hionine I:g14269] [GN:CO DE:Bacil	29928] nine uroporphyrinogen 5] [LN:BACCOBA] BA] [OR:Bacillus lus megaterium

 ORF Name
 NTID
 AAID
 NT AA Length Length
 Score
 probability

 A17503000985_38876_f2_481
 1196
 4968
 147
 48

 Description NO-HIT
 NO-HIT
 AAID
 AAID
 AAID
 Length Length
 48

ORF Name	NTID	AAID	NT AA score probability
A17503000985_3906718_£1_67	1197	4969	1065 354 747 5.2e-74
Description pir:[LN:H69789] [AC:H69789] [PN:L-i dehydrogenase homolog] [GN:ydjL] [C dehydrogenase homology] [OR:Bacillus >gp:[GI:d1023636:g2522016] [LN:AB007 [OR:Bacillus subtilis] [SR:Bacillus [DB:genpept-bct1] [DE:Bacillus subti 48degree.] [NT:putative] [LE:9403] [>gp:[GI:e1182603:g2632937] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-b 4 of 21): from 600701 to813890.] [NT [RE:77968] [DI:complement]	L:alcoho subtil: 638] [Ao subtili: lis geno RE:1044: 004] [Ao ct1] [D]	ol dehyis] [EGC: ABOO'S (straomic Di 3] [DI C: Z991(E: Baci	nydrogenase:long-chain alcohol C:1.1.1.14] [DB:pir1] 7638] [PN:dehydrogenase] [GN:ydjL] rain:Marburg 168) DNA] DNA containing gutA to cotA region, c:complement] .07:AL009126] [GN:ydjL] [FN:unknown] .1lus subtilis complete genome (section
ORF Name	NTID	AAID	NT AA core probability
A17503000985_3909376_c2_1681	1198	4970	396 131 244 1.0e-20
cassette homology] [OR:Helicobacter [AC:AE000655:AE000511] [PN:ABC trans	pylori] porter, enpept-l	[DB:pi ATP-bi bct2]	[DE:Helicobacter pylori 26695 section
ORF Name	NTID	AAID	NT AA score probability
A17503000985_3910675_£3_1222	1199	4971	234 77 57 0.0048
enzyme] [OR:Mitochondrion Aphidius e [DE:Aphidius ervi NADH dehydrogenase protein, partial cds.] [LE:<1] [RE:> [AC:AF069163] [PN:NADH dehydrogenase Aphidius picipes] [SR:Aphidius picip dehydrogenase 1 gene, mitochondrial [LE:<1] [RE:>465] [DI:direct] >gp:[G dehydrogenase 1] [FN:respiratory enz [SR:Aphidius pisivorus] [DB:genpept-	rvi] [SI 1 gene, 465] [DI 1] [FN: es] [DB: geneenco I:g32934 yme] [OI inv1] [I ochondri 069166] drion Ar i NADH o	R:Aphic , mitoo I:direc :respin :genper oding m 460] [I R:Mitoo DE:Aphi ial pro [AC:AI ohidius dehydro	chondrial geneencoding mitochondrial ct] >gp:[GI:g3293458] [LN:AF069163] ratory enzyme] [OR:Mitochondrion pt-inv1] [DE:Aphidius picipes NADH mitochondrial protein, partial cds.] LN:AF069164] [AC:AF069164] [PN:NADH chondrion Aphidius pisivorus] didius pisivorus NADH dehydrogenase 1 otein, partial cds.] [LE:<1] [RE:>465] F069166] [PN:NADH dehydrogenase 1] s sonchi] [SR:Aphidius sonchi] ogenase 1 gene, mitochondrial
ORF Name	NTID	AAID	NT AA score probability
A17503000985_3923842_c2_1832	1200	4972	<u>Length Length</u> <u>876</u> 1.1e-87
Description			J
<pre>gp:[GI:g4959402] [LN:AF115391] [AC:A [GN:rbsU] [OR:Lactobacillus sakei] [1 (laaA) gene, partial cds; LaaB (laaB genes, completecds; rbs operon, completects;</pre>	DB:genpe),putati	ept-bct ive ace	t2] [DE:Lactobacillus sakei LaaA etate kinase AckA (ackA), LaaC (laaC)

320

[LE:3130] [RE:4014] [DI:direct]

ORF Name	NTID	AAID Length Length score probability
A17503000985_3928162_f2_665	1201	4973 [141] 46
Description	الــــا	
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000985_3928416_f1_170	1202	4974 123 40
Description NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000985_3933177_£2_683	1203	4975 126 41
Description		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000985_3941078_f1_114	1204	4976 324 107 110 1.6e-06
Description		al protein yjbS] [GN:yjbS] [OR:Bacillus
<pre>subtilis] [DB:pir2] >gp:[GI:e1183188 [GN:yjbS] [FN:unknown] [OR:Bacillus</pre>	3:g26335 subtili	[LN:BSUB0007] [AC:Z99110:AL009126] s] [DB:genpept-bct1] [DE:Bacillus subtilis 4391to 1411140.] [LE:49925] [RE:50125]
		NT AA
ORF Name	NTID	AAID LengthLength score probability
ORF Name A17503000985_3941436_f1_6	NTID 1205	$\Delta\Delta III$ — — ecore probability
	-	LengthLength score probability
A17503000985_3941436_f1_6	-	LengthLength score probability
A17503000985_3941436_f1_6 Description	-	AAID LengthLength score probability AAID NT AA score probability
Description NO-HIT	1205	LengthLength score probability 4977 150 49 NT AA
Description NO-HIT ORF Name	NTID	AAID LengthLength score probability 4977 150 49 AAID NT AA LengthLength score probability
Description NO-HIT ORF Name AI7503000985_3953400_f1_314 Description sp:[LN:RFA1_KLEPN] [AC:Q48475] [GN:FEXPORT SYSTEM PERMEASE PROTEIN RFBA] [AC:S60882] [PN:integral membrane C[CL:integral membrane O-antigen translocator] [GN:wzm] [OR:Klebsiell pneumoniae integral membrane O-antig	NTID 1206 RFBA] [O: [SP:Q4- 0-antige: aslocato: FBA] [AC a pneumore gen transid WbbM	AAID LengthLength AAID NT AA LengthLength score probability AAID LengthLength score probability 4978 195 64 49 0.017 R:KLEBSIELLA PNEUMONIAE] [DE:O-ANTIGEN 8475] [DB:swissprot] >pir:[LN:S60882] at translocator protein rfbA] [GN:rfbA] r protein rfbA] [OR:Klebsiella pneumoniae] :L31775] [PN:integral membrane O-antigen oniae] [DB:genpept-bct2] [DE:Klebsiella slocatorprotein (wzm) and ATP-binding (wbbM) gene, partial cds.] [NT:similar to
Description NO-HIT ORF Name A17503000985_3953400_f1_314 Description sp:[LN:RFA1_KLEPN] [AC:Q48475] [GN:FEXPORT SYSTEM PERMEASE PROTEIN RFBA] [AC:S60882] [PN:integral membrane C[CL:integral membrane O-antigen translocator] [GN:wzm] [OR:Klebsiell pneumoniae integral membrane O-antigen translocat	NTID 1206 RFBA] [O: [SP:Q4- 0-antige: aslocato: FBA] [AC. a pneumore gen transid WbbM ,] [LE:: NTID	LengthLength AAID LengthLength AAID LengthLength LengthLength Score probability AAID LengthLength AP 0.017 R:KLEBSIELLA PNEUMONIAE] [DE:O-ANTIGEN B475] [DB:swissprot] >pir:[LN:S60882] A translocator protein rfbA] [GN:rfbA] Ar protein rfbA] [OR:Klebsiella pneumoniae] CL31775] [PN:integral membrane O-antigen CDiae] [DB:genpept-bct2] [DE:Klebsiella CSIOCATORYPOTE (WZM) and ATP-binding (wbbM) gene, partial cds.] [NT:similar to CS24] [RE:1103] [DI:direct] AAID NT AA LengthLength score probability
Description NO-HIT ORF Name AI7503000985_3953400_f1_314 Description sp:[LN:RFA1_KLEPN] [AC:Q48475] [GN:FEXPORT SYSTEM PERMEASE PROTEIN RFBA] [AC:S60882] [PN:integral membrane Cle:integral membrane O-antigen translocator] [GI:g567182] [LN:KPNRFtranslocator] [GN:wzm] [OR:Klebsiell pneumoniae integral membrane O-antigen translocator] [GN:wzm] [OR:wzm] [OR:wz	NTID 1206 RFBA] [O: [SP:Q4- 0-antige: aslocato: FBA] [AC. a pneumore gen transid WbbM ,] [LE:	LengthLength AAID LengthLength AAID LengthLength LengthLength score probability AP78 195 64 49 0.017 R:KLEBSIELLA PNEUMONIAE] [DE:O-ANTIGEN 8475] [DB:swissprot] >pir:[LN:S60882] a translocator protein rfbA] [GN:rfbA] r protein rfbA] [OR:Klebsiella pneumoniae] :L31775] [PN:integral membrane O-antigen pniae] [DB:genpept-bct2] [DE:Klebsiella slocatorprotein (wzm) and ATP-binding (wbbM) gene, partial cds.] [NT:similar to 324] [RE:1103] [DI:direct]

ORF Name	NTID AAID NT AA score probability
A17503000985 4069643 c3_2243	1208 4980 480 159 160 8.2e-12
Description	
gp:[GI:e1310302:g3294247] [LN:SC7C7] [GN:SC7C7.14] [OR:Streptomyces coeli	[AC:AL031031] [PN:hypothetical protein SC7C7.14] color] [DB:genpept-bct1] [DE:Streptomyces , unknown, len: 161 aa] [LE:24125] [RE:24610]
ORF Name	NTID AAID NT AA score probability
AI7503000985_4079552_f2_696	1209 4981 630 209 87 0.0012
[SR:Hepatitis A virus LSH/S] [DB:gen	4069] [GN:VP4/VP2] [OR:Hepatitis A virus] upept-vrl] [DE:VP13C/3D [hepatitis A virus HAV, uent 2 of 7].] [NT:This sequence comes from Fig. 3.]
ORF Name	NTID AAID NT AA score probability
AI7503000985_4079626_±1_336	1210 4982 192 63 70 0.018
Description	
[PN:serine proteinase inhibitor pred	rissprot] >gp:[GI:g438481] [LN:BOVSPIS] [AC:L22095] rursor] [OR:Bos taurus] [DB:genpept-mam] [DE:Bos or, mRNA, complete cds.] [NT:uterine milk protein]
ORF Name	NTID AAID NT AA score probability
A17503000985_4084635_f2_483	1211 4983 141 46
Description NO-HIT	
ORF Name	NTID AAID NT AA score probability
AI7503000985_4084717_f3_961	1212 4984 621 206 110 0.00027
<pre>subtilis] [DB:pir2] >gp:[GI:e1186182 [GN:yvpB] [FN:unknown] [OR:Bacillus complete genome (section 18 of 21): [DI:direct] >gp:[GI:g2618863] [LN:AF</pre>	othetical protein yvpB] [GN:yvpB] [OR:Bacillus :g2636007] [LN:BSUB0018] [AC:Z99121:AL009126] subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis from 3399551to 3609060.] [LE:189121] [RE:189873] [O17113] [AC:AF017113] [PN:YvpB] [GN:yvpB] ct2] [DE:Bacillus subtilis 300-304 degree genomic :complement]
ORF Name AI7503000985_4093932_f1_69 Description	NTID AAID NT AA score probability 1213 4985 135 44

ORF Name	NTID	AAID	Length Length score	probability
A17503000985_4094703_c1_1446	1214	4986	2022 673 224	3.7e-21
Description	L			
pir:[LN:B70001] [AC:B70001] [PN:ABC [OR:Bacillus subtilis] [DB:pir2] >gp [AC:Z99119:AL009126] [GN:ytsD] [FN:ur [DE:Bacillus subtilis complete genome [NT:similar to ABC transporter (perme >gp:[GI:g2293178] [LN:AF008220] [AC:A subtilis] [DB:genpept-bct2] [DE:Bacil [NT:similarity to NADH dehydrogenase]	:[GI:e1] nknown] e (sect: ease)] AF008220	185910 [OR:B ion 16 [LE:11 D] [PN	:g2635521] [LN:BSU acillus subtilis] of 21): from 2997 0708] [RE:112648] :YtsD] [GN:ytsD] [rrnB-dnaB genomic	[DB:genpept-bct1] 771to 3213410.] [DI:complement] OR:Bacillus region.]
ORF Name	NTID	AAID	NT AA score	probability
A17503000985_40966_c3_2016	1215	4987	1626 541	
Description NO-HIT	L			
ORF Name	NTID	AAID	NT AA LengthLength score	probability
A17503000985_4098518_£1_48	1216	4988	711 236 121	5.8e-08
sp:[LN:ARP_PLAFA] [AC:P04931] [OR:PLAFA] (AG319) (ARP) (FRAGMENT)] [SP:P04931] [PN:asparagine-rich protein] [OR:Plass [LN:PFAARP] [AC:M24328:X03716] [PN:asparagine] [SR:Plass falciparum] [SR:Plass falciparum] [DB:genpept-inv1] [DE:Plass falciparum] [DB:genpept-inv1] [DE:Plass falciparum] [LE:1] [RE:1612] [DI:direct]	[DB:sv smodium sparagir n (Ghana	visspr falci ne-ric nian i	ot] >pir:[LN:A2377 parum] [DB:pir2] > h protein] [GN:Ag3 solate NF7) asexua	0] [AC:A23770] gp:[GI:g160092] 19] [OR:Plasmodium l bloodstage]
ORF Name	NTID	AAID	NT AA score	probability
A17503000985_4100453_c3_2260	1217	4989	1044 347 329	1.0e-29
Description		L		
sp:[LN:DEGS_BACBR] [AC:P54663] [GN:DE PROTEIN DEGS,] [SP:P54663] [DB:swisspkinase] [GN:degS] [CL:regulatory pro>gp:[GI:g710495] [LN:BACDEGSU] [AC:L1 [OR:Brevibacillus brevis] [SR:Bacillu [DE:Bacillus brevis protein kinase (degu) gene, complete cds.]	orot] >rotein de 15444] 1s brevi degS) ge	oir:[Li egS] [0 [PN:pro s (st: ene, co	N:I39834] [AC:I398 OR:Bacillus brevis otein kinase] [GN: rain Alk36) DNA] [I omplete cds;transc] [DB:pir2] degS] DB:genpept-bct2] riptional activator
ORF Name	NTID	AAID	NT AA score	probability
A17503000985_4100938_c2_1860	1218	4990	534 177 293	1.1e-25
Description				
sp:[LN:GUDT_BACSU] [AC:P42237] [GN:YCTRANSPORTER] [SP:P42237] [DB:swissproglucarate transporter] [GN:ycbE] [CI[DB:pir2] >gp:[GI:d1007040:g709999] [dehydratase] [GN:ycbE] [OR:Bacillus schromosomecontaining yckA-T genes.] [>gp:[GI:e1182200:g2632534] [LN:BSUB00]	ot] >pir ::hexurc [LN:BACY subtilis subtilis [LE:3924	c:[LN:] cnate (CB20] c] [SR c DNA c	H69752] [AC:H69752 transporter] [OR:Ba [AC:D30808] [PN:gl :Bacillus subtilis around 20 degrees n :5291] [DI:direct]] [PN:probalble acillus subtilis] lucarate (strain:168TrpC2) region of

ORF Name	NTID	AAID LengthLength score probability
A17503000985_4110712_c1_1353	1219	4991 591 196
Description	<u> </u>	
NO-HIT		
		NITT A A
ORF Name	NTID	AAID LengthLength score probability
A17503000985_4112527_f2_563	1220	4992 144 47
Description	<u> </u>	
NO-HIT		
		NT AA
ORF Name	NTID	AAID LengthLength score probability
A17503000985_4112788_c1_1497	1221	4993 141 46
Description		
NO-HIT		
Control of the Contro		NT AA
ORF Name	NTID	AAID LengthLength score probability
AI7503000985_4148428_f1_160	1222	4994 1149 382 528 8.3e-51
Description		
sp:[LN:ADH_ALCEU] [AC:P14940] [GN:AD		- _ -
		:swissprot] >pir:[LN:A30196] [AC:A30196]
		cohol dehydrogenase:long-chain alcohol
dehydrogenase homology] [OR:Alcalige		R:Ralstonia eutropha] [SR:A.eutrophus
		ctl] [DE:A.eutrophus alcohol dehydrogenase
		drogenase (EC 1.1.1.1)] [LE:458] [RE:1558]
[DI:direct]		
ODD Ware		NT AA
ORF Name	NTID	AAID LengthLength score probability
AI7503000985_4164026_f2_793	1223	4995 162 53 118 9.1e-09
Description		
pir:[LN:S68609] [AC:S68609] [PN:rec		
[OR:Staphylococcus aureus] [DB:pir2]		
		s aureus] [SR:Staphylococcus aureus DNA; :genpept-bct1] [DE:Staphylococcus aureus
	פתו נחם	
	ויידא] וו	outativel [LE-389] [RE-997] [DI-direct]
	[NT:]	putative] [LE:389] [RE:997] [DI:direct]
ORF Name	NTID	NIT AA
	NTID	AAID NT AA score probability
AI7503000985_4167842_f2_530		NIT AA
	NTID	AAID NT AA score probability
A17503000985_4167842_f2_530 Description	NTID	AAID NT AA score probability LengthLength [4996] 201 66
A17503000985_4167842_f2_530 Description	NTID	AAID NT AA score probability AAID NT AA score probability
AI7503000985_4167842_f2_530 Description NO-HIT ORF Name	NTID 1224 NTID	AAID NT AA score probability 4996 201 66 AAID NT AA probability AAID NT AA probability Length Length Score probability
AI7503000985_4167842_f2_530 Description NO-HIT	NTID 1224	AAID NT AA score probability AAID NT AA score probability

ORF Name	NTID AAID NT AA score probability
A17503000985 4303377_t2_610] [1226] [4998] [270] [89] [81] [0.0019
Description	
gp:[GI:g4049784] [LN:AF063866] [AC:A [GN:MSV234] [OR:Melanoplus sanguining	AF063866] [PN:ORF MSV234 hypthetical protein] pes entomopoxvirus] [DB:genpept-vrl] [DE:Melanoplus e genome.] [LE:201477] [RE:201830] [DI:complement]
ORF Name	$rac{ ext{NTID}}{ ext{Length}} = rac{ ext{NT}}{ ext{Length}} rac{ ext{Score}}{ ext{probability}}$
A17503000985_4329453_c1_1533 Description	1227 4999 336 111 220 3.6e-18
subtilis] [DB:pir2] >gp:[GI:d1020138] [OR:Bacillus subtilis] [SR:Bacillus [DE:Bacillus subtilis genome sequence degree.] [NT:SIMILAR TO THIOREDOXIN. >gp:[GI:e1182517:g2632851] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-b	ioredoxin homolog ydfQ] [GN:ydfQ] [OR:Bacillus 8:g1881358] [LN:AB001488] [AC:AB001488] [GN:ydfQ] subtilis (strain:168) DNA] [DB:genpept-bct1] ce, 148 kb sequence of the regionbetween 35 and 47 .] [LE:131947] [RE:132285] [DI:direct] 0003] [AC:Z99106:AL009126] [GN:ydfQ] [FN:unknown] bct1] [DE:Bacillus subtilis complete genome (section T:similar to thioredoxin] [LE:195693] [RE:196031]
ORF Name	NTID AAID NT AA score probability
A17503000985_4334818_£2_854	1228 5000 1566 521 111 0.018
[OR:Plasmodium falciparum] [DB:pir2] [AC:AE001385:AE001362] [PN:hypothetifalciparum] [SR:malaria parasite P.	ical protein] [GN:PFB0285c] [OR:Plasmodium falciparum] [DB:genpept-inv2] [DE:Plasmodium of 73 of thecomplete sequence.] [NT:predicted by
ORF Name	NTID AAID NT AA score probability
AI7503000985_4336088_c3_2078	1229 5001 624 207 283 7.6e-25
[GN:ylnF] [CL:Aquifex aeolicus sirc >gp:[GI:e332189:g2462962] [LN:BSPYRE [OR:Bacillus subtilis] [DB:genpept-bregion.] [LE:5769] [RE:6257] [DI:dir [AC:Z99112:AL009126] [GN:ylnF] [FN:u [DE:Bacillus subtilis complete genom	oporphyrin-III C-methyltransferase homolog ylnF] oheme synthase] [OR:Bacillus subtilis] [DB:pir2] EYLO] [AC:AJ000974] [PN:YlnF protein] [GN:ylnF] oct1] [DE:Bacillus subtilis pyrE to yloA gene rect] >gp:[GI:e1185155:g2633936] [LN:BSUB0009] unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] me (section 9 of 21): from 1598421to 1807200.] ethyltransferase] [LE:36582] [RE:37070] [DI:direct]
ORF Name A17503000985_4379763_f3_1048 Description	NTID AAID NT AA score probability 1230 5002 123 40

ORF Name	NTID	AAID	<u>NT</u> Length	AA Length so	core	probability
AI7503000985_4392767_c1_1619	1231	5003	435	144	96 8	.0e-15
Description				``		
<pre>gp:[GI:e1314293:g3395543] [LN:SC4A2] [GN:SC4A2.10c] [OR:Streptomyces coel coelicolor cosmid 4A2.] [NT:SC4A2.10 [RE:11015] [DI:complement]</pre>	icolor]	[DB:ge	enpept-	bct1] [I	E:Str	eptomyces
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	core	probability
AI7503000985_4459375_£2_830	1232	5004		68		
Description NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	core	probability
A17503000985_4487550_f3_980	1233	5005	1707	568 1	408 4	.7e-144
Description						
pir:[LN:D69748] [AC:D69748:I40419] [OR:Bacillus subtilis] [DB:pir2] >gp [GN:ybeC] [OR:Bacillus subtilis] [SR [DB:genpept-bct1] [DE:Bacillus subti 23degree.] [LE:34338] [RE:35957] [DI [AC:Z99105:AL009126] [GN:ybeC] [FN:u [DE:Bacillus subtilis complete genom [NT:similar to amino acid transporte	:[GI:d10 :Bacillu lis geno :direct] nknown] e (sect:	034075: us subt omic DN] >gp: [OR:Ba ion 2 c	:g35996 cilis (NA, 70 [GI:e11 acillus of 21):	34] [LN: strain:1 kb regio 82164:g2 subtili from 19	AB006 .68) D on bet .63249 .s] [D	424] [AC:AB006424] NA] ween 17 and 8] [LN:BSUB0002] B:genpept-bct1] to415810.]
ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	core	probability
A17503000985_4491713_f2_533 Description	1234	5006				.1e-39
pir:[LN:F64554] [AC:F64554] [PN:gua [OR:Helicobacter pylori] [DB:pir2] > [AC:AE000546:AE000511] [PN:guanosine [OR:Helicobacter pylori 26695] [DB:g 24 of 134 of the complete genome.] [PID:148183] [LE:6964] [RE:8418] [DI:	gp:[GI:g pentapl enpept-l NT:simi]	g231336 nosphat oct2]	8] [LN e phos [DE:Hel	:AE00054 phohydro icobacte	6] lase] r pylo	[GN:HP0278] ori 26695 section
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length sc	ore	probability
AI7503000985_4553166_£3_1122	1235	5007		<u>57</u>		
Description NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	AA Length	ore	probability
AI7503000985_4574012_f1_208	1236	5008			00 2.	.0e-58
Description						
<pre>pir:[LN:JN0500] [AC:JN0500:S34967] mitochondrial:dihydrocrotate_dehydroc</pre>	_				roorot	tate oxidadel

[OR:Drosophila melanogaster] [EC:1.3.3.1] [DB:pir2]

ORF Name	NTID	AAID LengthLength score probability
A17503000985_4662_t2_689	1237	5009 318 105 84 10.0046
Description	L	
		PN:ORF MSV064 hypothetical protein
		mopoxvirus] [DB:genpept-vrl] [DE:Melanoplus .] [LE:64316] [RE:65158] [DI:complement]
sanguinipes encompositius, complete	genome	.] [DE:04310] [OCICO:AN] [OCICO:AN]
ORF Name	NTID	AAID NT AA score probability LengthLength
AI7503000985_4687705_c2_1841	1238	5010 141 46
Description		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000985_4687752_c1_1514	1239	5011 1185 394 773 9.1e-77
Description		
-		aminotransferase homolog ykrV] [GN:ykrV]
-		otilis] [DB:pir2] >gp:[GI:e1184948:g2633729] [FN:unknown] [OR:Bacillus subtilis]
	_	plete genome (section 8 of 21): from
	spartate	e aminotransferase] [LE:30317] [RE:31513]
[DI:direct]		
ORF Name	NTID	AAID <u>NT AA</u> score probability
A17503000985_4689390_c3_2124	1240	5012 147 48
Description		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000985_4694163_c2_1791	1241	5013 144 47
Description		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000985_4694652_c3_2199	1242	5014 135 44
Description		
NO-HIT		
		AND NT AA ggore probability
ORF Name	<u>NTID</u>	AAID LengthLength score probability
A17503000985_4695451_f2_864	1243	5015 135 44
Description		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000985_4698428_c2_1705	1244	5016 180 59
Description		
NO-HIT		

ORF Name A17503000985_4705077_f2_880	NTID 1245	AAID NT AA Score probability 5017 123 40
Description NO-HIT		
ORF Name A17503000985_4713377_f2_618	NTID 1246	AAID NT AA score probability [5018 447 148 108 3.4e-05
	falcipar	
ORF Name AI7503000985_4719775_c3_2182	NTID 1247	AAID NT AA score probability 5019 744 247 434 7.6e-41
24.6 KD PROTEIN IN DAE-TYRZ INTERGEN >pir: [LN:S39662] [AC:S39662:C70050] ipa-7d] [GN:ywaC] [CL:GTP pyrophosp [DB:pir2] >gp:[GI:g413931] [LN:BSGEN [DB:genpept-bct1] [DE:B.subtilis gen [RE:7613] [DI:direct] >gp:[GI:e11863 [GN:ywaC] [FN:unknown] [OR:Bacillus	IC REGIO [PN:GTF hokinase R] [AC:X omic reg 47:g2636 subtilis from 379	P-pyrophosphokinase homolog ywaC:protein e related protein] [OR:Bacillus subtilis] [K73124] [GN:ipa-7d] [OR:Bacillus subtilis] [GN:ipa-7d] [SP:P39583] [LE:6981] [GN:BSUB0020] [AC:Z99123:AL009126] [DB:genpept-bct1] [DE:Bacillus subtilis [B8401to 4010550.] [NT:alternate gene name:
ORF Name A17503000985_4719827_c3_1986 Description NO-HIT	NTID 1248	AAID NT AA score probability [5020] 417 [138]
aureus] [DB:genpept-bct2] [DE:Staphy	lococcus	AAID NT AA LengthLength score probability [5021 1257 418 1659 1.2e-170 [PN:FmhA] [GN:fmhA] [OR:Staphylococcus aureus FmhA (fmhA) gene, complete cds.] ad FemB] [LE:201] [RE:1451] [DI:direct]
ORF Name A17503000985_4723510_c2_1759 Description	NTID 1250	AAID NT AA score probability LengthLength 2.0e-246

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sp:[LN:PODK_CLOSY] [AC:P22983] [GN:PPDK] [OR:CLOSTRIDIUM SYMBIOSUM] [SR:,BACTEROIDES
SYMBIOSUS] [EC:2.7.9.1] [DE:DIKINASE)] [SP:P22983] [DB:swissprot]

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Lengtl	score	probability
AI7503000985_4725068_c3_2219	1251	5023	645	214	605	5.8e-59
Description						
pir:[LN:D69670] [AC:D69670] [PN:gly (membrane p) opuCB] [GN:opuCB] [CL: [OR:Bacillus subtilis] [DB:pir2] >gp [PN:transmembrane protein] [GN:opuCB [DE:Bacillus subtilis osmoprotectant transmembrane protein (opuCB), osmop transmembrane protein (opuCD)genes, osmoprotectant transport system] [LE >gp:[GI:e1186070:g2635895] [LN:BSUBO betaine/carnitine/choline ABC] [GN:obetaine,] [OR:Bacillus subtilis] [DB genome (section 18 of 21): from 3399 [LE:68697] [RE:69350] [DI:complement	glycine :[GI:g2:] [OR:Batransparentectar complete:2025] 018] [AapuCB] [Senpep:	betain 271390 acillumont synthind e cds. [RE:26 C:2991 FN:high	ne/carne/carne s subt: stem Oping pro [NT:078] [D: 78] [D:21:ALO06 h affin	nitine AF0093 ilis] puC in otein OpuCB; I:dire 09126] nity t Bacill	/cholin 52] [A0 [DB:ger cluding precurs part o ct] [PN:g] ranspon us subt	ne ABC transporter] C:AF009352] npept-bct1] gATPase (opuCA), sor (opuCC) and of the Lycine ct of glycine cilis complete
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Lengtl	score	probability
A17503000985_4726636_c1_1543	1252	5024	138	45	7	
Description		<u> </u>	- ,		_	
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Lengtl	score	probability
AI7503000985_4727187_c2_1702	1253	5025	2889	962	617	7.4e-58
sp:[LN:PIP_LACLA] [AC:P49022] [GN:PI [SR:,SUBSPLACTIS:STREPTOCOCCUS LACTI [DB:swissprot] >gp:[GI:g308861] [LN:infection] [OR:Lactococcus lactis] [lactis) DNA] [DB:genpept-bct1] [DE:Lcds's, and rrggene, 5' end of cds.]	S] [DE:1 LACPIP] SR:Lacto actococo	PHAGE : [AC:L: ococcus cus lac	INFECT: 14679] s lact: ctis p:	ION PRO [GN:p: is (st: ip and	OTEIN] ip] [Fi rain C2 gerC2	V:required for phage 2, sub_species genes, complete
ORF Name	NTID	AAID	NT Length	AA Length	score	probability
A17503000985_4728187_c1_1365	1254	5026	234	77	<u>.</u>]	
Description NO-HIT					_	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Lengtl	score	probability
AI7503000985_4735937_f3_1114	1255	5027	138	45	7	
Description			-,		J	
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000985_4737550_c3_2248	1256	5028	759	252	387	7.3e-36
Description						
<pre>gp:[GI:g4433637] [LN:AF029224] [AC:A in nitrite reduction] [OR:Staphyloco carnosus nir and nar operons, comple</pre>	ccus car	rnosus]	[DB:g	genpept	-bct2]	[DE:Staphylococcus

ORF Name	NTID	AAID	NT AA core probability
AI7503000985_4777217_c3_2257	1257	5029	
product] [LE:7804] [RE:9381] [DI:dir [AC:AF029224:AF029225] [PN:NarH] [GN	nce 1 f ect] >g :narH] carnos	rom Pa p:[GI: [OR:St us nir	atent EP0805205.] [NT:unnamed protein:g3929523] [LN:AF029224] taphylococcus carnosus] r and nar operons, complete sequences.]
ORF Name	NTID	AAID	NT AA LengthLength
AI7503000985_4787807_£3_984	1258	5030	
Description NO-HIT			
ORF Name	NTID	AAID	NT AA LengthLength score probability
AI7503000985_4814682_£3_1288	1259	5031	201 66
Description NO-HIT			
ORF Name	NTID	AAID	NT AA LengthLength score probability
AI7503000985_4870907_£1_195	1260	5032	<u>5</u> <u> </u>
Description NO-HIT		•	
ORF Name	NTID	AAID	NT AA LengthLength
AI7503000985_4875316_c1_1545	1261	5033	141 46
Description NO-HIT			
ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000985_4875760_f1_255	1262	5034	135 44
Description NO-HIT	and the same of th		
ORF Name	NTID	AAID	NT AA Probability
A17503000985_4876387_c3_2099	1263	5035	186 61
Description NO-HIT			
ORF Name	NTID	AAID	NT AA LengthLength score probability
AI7503000985_4876932_c2_1733	1264	5036	249 82 74 0.013
Description			
<pre>gp:[GI:g1255424] [LN:CELC33G8] [AC:U [DB:genpept-inv1] [DE:Caenorhabditis [RE:27848:27952:28290] [DI:directJoin</pre>	elegans		

ORF Name	NTID	AAID LengthLength score probability
A17503000985_4878312_c2_1648	1265	5037 141 46
Description		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000985_4882893_c3_2012	1266	5038 1209 402 244 1.0e-27
Description		
<pre>gp:[GI:d1039027:g4512424] [LN:AB0175 halodurans] [SR:Bacillus halodurans halodurans C-125 genomic DNA, 32 kb (identity of 70% to B. subtilis)] [</pre>	(strain fragmen	:C-125) DNA] [DB:genpept-bct1] [DE:Bacillus at, completecds.] [NT:secY homologue
ORF Name	NTID	AAID NT AA roote probability
A17503000985_4891002_c2_1834	1267	5039 1116 371 671 5.8e-66
Description		
pir:[LN:A43577] [AC:A43577] [PN:reg [DB:pir2]	gulatory	protein pfoR] [OR:Clostridium perfringens]
ORF Name	NTID	AAID NT AA score probability
AT7503000985_4980378_c1_1635	1268	5040 537 178 578 4.2e-56
Description		
<pre>gp:[GI:g4096799] [LN:SCU40158] [AC:U [DB:genpept-bct2] [DE:Staphylococcus (orfx)gene, partial cds.] [NT:orfx; [RE:560] [DI:direct]</pre>	carnos	
ORF Name	NTID	AAID NT AA score probability
A17503000985_5117137_c2_1827	1269	5041 459 152 499 9.8e-48
Description		
[GN:ywtA] [OR:Bacillus subtilis] [DE:Z99122:AL009126] [GN:ywtA] [FN:u] [DE:Bacillus subtilis complete genom [NT:similar to capsular polyglutamat [DI:complement] >gp:[GI:e308089:g189] [OR:Bacillus subtilis] [DB:genpept-bgenes.] [NT:product highly similar to [DI:direct] >gp:[GI:e1184495:g263611] [FN:unknown] [OR:Bacillus subtilis]	DB:pir2] unknown] ne (sect: e biosyn 4766] [1 cct1] [DI co Bacil: 4] [LN:I [DB:gen 809700.]	LN:BSZ92954] [AC:Z92954] [GN:ywtA] E:B.subtilis yws[A,B,C,D,E,F,G] and gerBC lus anthracis CapC] [LE:1084] [RE:1533] BSUB0019] [AC:Z99122:AL009126] [GN:ywtA] pept] [DE:Bacillus subtilis complete genome] [NT:similar to capsular polyglutamate
ORF Name	NTID	AAID NT AA score probability
AI7503000985_5128425_£2_821	1270	5042 1878 625 477 2.1e-45
Description		
[GN:yjdC] [CL:phosphotransferase sy homology] [OR:Bacillus subtilis] [DB [AC:Z99110:AL009126] [GN:yjdC] [FN:u [DE:Bacillus subtilis complete genom	stem mar :pir2] : nknown] e (secti	ion antiterminator BglG family homolog yjdC] nnitol-specific enzyme II factor III >gp:[GI:e1183220:g2633554] [LN:BSUB0007] [OR:Bacillus subtilis] [DB:genpept-bct1] ion 7 of 21): from 1194391to 1411140.] or (BglG] [LE:75712] [RE:77658] [DI:direct]

ORF Name	NTID	AAID	<u>NT</u> Lengtl	<u>AA</u> hLength	score	probability
A17503000985_5133562_£2_656	1271	5043	222	73	180	6.3e-14
Description						
pir:[LN:F70041] [AC:F70041] [PN:promercuric ion-binding protein merP horegulatory protein: heavy-metal-assocyp:[GI:e1186039:g2635864] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-hammed heavy from 3399551to 3609060.] [LE:43125] [RE:43334] [DI:complement	omolog] ociated 0018] [A octl] [D [NT:sim	[GN:yv homolo C:Z991 E:Baci	gy] [gy] [O 21:AL0 llus s	CL:mero R:Baci 09126] ubtilis	curic : llus su [GN:yv s comp]	resistance operon ubtilis] [DB:pir1] vgY] [FN:unknown] Lete genome (section
ORF Name	NTID	AAID	NT Lengtl	<u>AA</u> hLength	score	probability
A17503000985_5136002_c2_1691	1272	5044	366	121		2.9e-09
Description	· ———			J L		
pir:[LN:C70043] [AC:C70043] [PN:hypsubtilis] [DB:pir2] >gp:[GI:e1186201 [GN:yvlA] [FN:unknown] [OR:Bacillus complete genome (section 18 of 21): [DI:complement] >gp:[GI:e1184419:g26 [FN:unknown] [OR:Bacillus subtilis] genome (section 19 of 21): from 3597 [DI:complement] >gp:[GI:g2618844] [ICOR:Bacillus subtilis] [DB:genpept-ksequence.] [LE:19550] [RE:19876] [DI:AC:Z99122:AL009126] [GN:yvlA] [FN:ucomplement] [DE:Bacillus subtilis complete genom [LE:10417] [RE:10743] [DI:complement]	l:g26360 subtili from 33 536039] [DB:gen 7091to 3 [N:AF017 oct2] [D: ct2] [D: ct2] [D: direct unknown]	26] [L s] [DB 99551t [LN:Bs pept-b 809700 113] [E:Baci] >gp: [OR:B	N:BSUB :genpe o 3609 UB0019 ct1] [LE .] [LE AC:AF0 llus s [GI:e1	0018] pt-bct1 060.]] [AC:2 DE:Baci :10417] 17113] ubtilis 184419: s subti	[AC: Z99 L] [DE: [LE: 207 Z99122: illus s [RE: 1 [PN: Yv s 300-3 :g26360 illis]	P121:AL009126] Bacillus subtilis P957] [RE:208283] AL009126] [GN:yvlA] Bubtilis complete D0743] VlA] [GN:yvlA] B04 degree genomic D39] [LN:BSUB0019] [DB:genpept]
ORF Name	NTID	AAID	<u>NT</u> Length	AA nLength	score	probability
A17503000985_5203452_c3_1989	1273	5045	579	192	398	5.0e-37
Description sp:[LN:HIS5_LACLA] [AC:Q02132] [GN:HIS5_LACLA] [AC:Q02132] [GN:HISR:,SUBSPLACTIS:STREPTOCOCCUS LACTI [SP:Q02132] [DB:swissprot] >pir:[LN:hisH] [OR:Lactococcus lactis subsp. [AC:U92974:M90760:M90761] [PN:HisH] [DE:Lactococcus lactis unknown gene, unknown, HisB (hisB), unknown, HisH unknown, unknown, LeuA(leuA), LeuB (IlvD(ilvD), IlvB (ilvB), IlvN, IlvC genes, complete cds.] [NT:aminotrans	[S] [EC:: :145734] lactis] [GN:his] partia: (hish),1 (leuB), 1 (ilvC),	2.4.2. [AC:I [DB:p h] [OR l cds, HisA (LeuC (IlvA	-] [DE 45734 ir2] >: :Lacto and H hisA), leuC), (ilvA)	:AMIDOT [PN:H gp:[GI: coccus isC (hi HisF (LeuD (, AldB	TRANSFE JISH] 1925651 1actis SC), un (hisF), (leuD), (aldB)	CL:amidotransferase [LN:LLU92974] [CB:genpept-bct1] [CR:Genpept-bct1] [CR:Genpept-bc
ORF Name A17503000985_5281568_f3_1043 Description	NTID 1274	<u>AAID</u> 5046	504	AA Length	330	probability 8.0e-30
pir:[LN:C64897] [AC:C64897] [PN:pro [CL:phosphinothricin N-acetyltransfe						

>gp:[GI:g1787719] [LN:AE000241] [AC:AE000241:U00096] [PN:putative resistance protein] [GN:b1448] [FN:putative transport; Drug/analog sensitivity] [OR:Escherichia coli] [DB:genpept-bct2] [DE:Escherichia coli K-12 MG1655 section 131 of 400 of the completegenome.] [NT:f172; 38 pct identical (3 gaps) to 169 residues of] [LE:9570] [RE:10088] [DI:complement]

AΑ NT ORF Name NTID AAID probability score LengthLength

AI7503000985 5290675 c3 2108

1275

5047 405

2.1e-38

Description

sp:[LN:PAND BACSU] [AC:P52999] [GN:PAND] [OR:BACILLUS SUBTILIS] [EC:4.1.1.11] [DE:DECARBOXYLASE)] [SP:P52999] [DB:swissprot] >pir:[LN:A69672] [AC:A69672] [PN:aspartate 1-decarboxylase panD] [GN:panD] [CL:aspartate 1-decarboxylase] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:g1146242] [LN:BACYPIA] [AC:L47709] [PN:aspartate 1-decarboxylase] [GN:panD] [FN:pantothenic acid biosynthesis] [OR:Bacillus subtilis] [DB:genpept-bct1] [EC:4.1.1.11] [DE:Bacillus subtilis (clone YAC15-6B) ypiABF genes, qcrABC genes, ypjABCDEFGHI genes, birA gene, panBCD genes, dinG gene, ypmB gene, aspB gene, asnS gene, dnaD gene, nth gene and ypoC gene, completecds's.] [NT:48.3% of identity to the Escherichia coli aspartate] [LE:14990] [RE:15373] [DI:direct] >gp:[GI:e1183686:g2634659] [LN:BSUB0012] [AC:Z99115:AL009126] [PN:aspartate 1-decarboxylase] [GN:panD] [FN:pantothenate biosynthesis] [OR:Bacillus subtilis] [DB:genpept-bct1] [EC:4.1.1.11] [DE:Bacillus subtilis complete genome (section 12 of 21): from 2195541to 2409220.] [SP:P52999] [LE:156294] [RE:156677] [DI:complement]

NT AA ORF Name NTID AAID score probability LengthLength A17503000985 5314077 c1 1448 5048 1593 1.2e-104

Description

sp:[LN:YIDK ECOLI] [AC:P31448] [GN:YIDK] [OR:ESCHERICHIA COLI] [DE:HYPOTHETICAL 62.1 KD PROTEIN IN EMRD-GLVG INTERGENIC REGION] [SP:P31448] [DB:swissprot] >pir:[LN:H65169] [AC:H65169] [PN:hypothetical 62.1 kD protein in ilvo-ibpb intergenic region] [GN:yidK [OR:Escherichia coli] [DB:pir2] >gp:[GI:g290528] [LN:ECOUW82] [AC:L10328] [GN:f571] [FN:unknown] [OR:Escherichia coli] [SR:Escherichia coli K12 strain MG1655; lambda clones EC14-52] [DB:genpept-bct1] [DE:E. coli; the region from 81.5 to 84.5 minutes.] [NT:similar to glucose transport proteins] [LE:47795] [RE:49510] [DI:complement] >gp:[GI:g1790113] [LN:AE000445] [AC:AE000445:U00096] [PN:putative cotransporter] [GN:yidK] [FN:putative transport; Not classified] [OR:Escherichia coli] [DB:genpept-bct2] [DE:Escherichia coli K-12 MG1655 section 335 of 400 of the completegenome.] [NT:f571; 100 pct identical to YIDK ECOLI SW:] [LE:5576] [RE:7291] [DI:complement]

NT AAORF Name NTID AAID <u>sco</u>re probability LengthLength A17503000985_53413_f2_855 1277 5049 1602 533 8.7e-40 424

Description

pir: [LN:H64532] [AC:H64532] [PN:2',3'-cyclic-nucleotide 2'-phosphodiesterase, precursor] [GN:HP0104] [CL:2',3'-cyclic-nucleotide 2'-phosphodiesterase: 2',3'-cyclic-nucleotide 2'-phosphodiesterase homology:phosphoesterase core homology] [OR:Helicobacter pylori] [EC:3.1.4.16] [DB:pir1] >qp:[GI:q2313187] [LN:AE000532] [AC:AE000532:AE000511] [PN:2',3'-cyclic-nucleotide 2'-phosphodiesterase] [GN:HP0104] [OR:Helicobacter pylori 26695] [DB:genpept-bct2] [DE:Helicobacter pylori 26695 section 10 of 134 of the complete genome.] [NT:similar to GB:L42023 SP:P44764 PID:1004048] [LE:3594] [RE:5339] [DI:complement]

ORF Name	NTID	AAID	NT Length Le	AA ength score	probability
AI7503000985_5369212_c3_2196	1278	5050	1509 5	02 430	5.4e-40
Description sp:[LN:TAGE_BACSU] [AC:P13484] [GN:TELEN:GEC 2.4.1.52) (TEICHOIC ACID BIG pir:[LN:S06048] [AC:S06048:F69720] alpha-glucosyltransferase, tagE:proken phosphate glucosyltransferase tagE] 2.4.1.52] [DB:pir2] [MP:310 degrees [OR:Bacillus subtilis] [DB:genpept-ken (gtaA) polypeptide (AA 1-673)] [SP:Fomology:[GI:e1184479:g2636099] [LN:BSUBG [PN:UDP-glucose:polyglycerol phosphaten [OR:Bacillus subtilis] [DB:genpept-ken (gtaA) gtaD] [SP:P13484] [LE:80369] [SP:P13484] [LE:80369] [SP:P13484] [LE:80369] [SP:P13484] [CR:Bacillus subtilis] [DB:genpept-ken (gtaA) gtaD] [SP:P13484] [LE:80369] [SP:P13484] [LE:80369] [SP:P13484] [LE:80369] [SP:P13484] [LE:80369] [SP:P13484] [LE:80369] [SP:P13484] [LE:80369] [RE:82390] [SP:P13484] [LE:80369] [RE:82390] [SP:P13484] [LE:80369] [RE:82390] [SP:P13484] [LE:80369] [RE:82390]	FAGE:ROD DSYNTHES [PN:po bable ro [GN:tag] >gp:[pct1] [D P13484] [O19] [A te] [GN pct1] [E 7091to 3 [RE:8239 [RE:8239 [GN] [GN [EC:2.4	D:GTAA IS PRO ly(gly dD pro E:rodD GI:g58 E:Baci [LE:15 C:Z991 :tagE] C:2.4. 809700 0] [DI C:Z991 :tagE] .1.52]] [NT:	[OR:BACTEIN E)] cerol-photein:UDP- [OR:BACTEIN E)] [OR:BACTEIN	CILLUS SUBT [SP:P13484 osphate) -glucosep acillus sub N:BSRODC] [tilis rodC 178] [DI:di 126] choic acid E:Bacillus lternate ge ent] 126] choic acid illus subti	ILIS] [EC:2.4.1.52]] [DB:swissprot] olyglycerol tilis] [EC: AC:X15200] operon.] [NT:rodD rect] biosynthesis] subtilis complete ne name: rodD, biosynthesis] lis complete genome
[attended] [attended] [attended]			- 46 ****	The state of the s	
ORF Name	NTID	AAID	NT Length Le	AA ength	probability
AI7503000985_54207_c1_1347	1279	5051	138 4	5	
Description NO-HIT					
ORF Name	NTID	AAID	NT LengthLe	AA ength	probability
AI7503000985_56693_c1_1418	1280	5052	792 20	63 778	2.7e-77
Description gp:[GI:g2058476] [LN:BTU71200] [AC:U [DB:genpept-mam] [DE:Bos taurus acet acetoin reductase of Klebsiella] [LE	oin red	uctase	mRNA, co	omplete cds	-
ORF Name	NTID	AAID	NT Length Le	AA ength score	probability
AI7503000985_578162_f3_1226 Description NO-HIT	1281	5053	177 58	В	
ORF Name	NTID	AAID	NT LengthLe	AA ength score	probability
AI7503000985_5867262_c1_1636	1282	5054	126 4		
Description NO-HIT		<u> </u>	JI L		
ORF Name AI7503000985_5891075_f2_736 Description	NTID 1283	<u>AAID</u> 5055	NT LengthLe		probability
NO-HIT					

ORF Name	NTID	AAID NT AA score probability
A17503000985_5907177_£1_392	1284	5056 138 45
Description		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000985_5915653_£2_789	1285	5057 183 60
Description		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000985_595260_c2_1956	1286	5058 3780 1259 5568 0.0
Description		
carnosus] [DB:genpept-pat] [DE:Seque product] [LE:4140] [RE:7814] [DI:dir [AC:AF029224:AF029225] [PN:NarG] [GN [DB:genpept-bct2] [DE:Staphylococcus	nce 1 f: ect] >gj :narG] carnosi	
ORF Name	NTID	AAID NT AA score probability
		Length Length Plobability
A17503000985_5963300_£2_895	1287	5059 504 167 116 5.6e-06
<pre>protein] [GN:APE1673] [OR:Aeropyrum</pre>	pernix]	AP000062] [PN:213aa long hypothetical [SR:Aeropyrum pernix (strain:K1) DNA] NA, section 5/7.] [LE:71354] [RE:71995]
ORF Name	NTID	AAID NT AA score probability
A17503000985_5988786_f1_391	1288	
Description		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000985_6047827_c2_1778	1289	5061 705 234 569 3.8e-55
Description		
cds, and HisC (hisC), unknown, HisG (hish), HisA (hisA), HisF (hisF), His (leuB), LeuC (leuC), LeuD (leuD), un IlvA (ilvA), AldB (aldB) and aldR (aldecarboxylase] [LE:24488] [RE:25198] [AC:S82499] [PN:alpha-acetolactate d [SR:Lactococcus lactis NCDO2118 ssp.	bctl] [IhisG), thisG), the hisG his land land land land land land land land	DE:Lactococcus lactis unknown gene, partial inknown, HisB (hisB), unknown, HisH (E), unknown, unknown, LeuA(leuA), LeuB (lvD(ilvD), IlvB (ilvB), IlvN, IlvC (ilvC), es, complete cds.] [NT:alpha-acetolactate eect] >gp:[GI:g1699352] [LN:S82499] (lase] [GN:aldB] [OR:Lactococcus lactis] [DB:genpept-bct1] (actococcus lactis, ssp.lactis, NCDO2118,
[DI:direct]		

ORF Name	NTID	AAID NT AA score probability
A17503000985_6070938_c1_1477	1290	
[OR:Staphylococcus xylosus] [DB:geng transporter (cudT), putativeregulate	ept-bct ory prot	[GN:cudB] [PN:choline dehydrogenase] [GN:cudB] [DE:Staphylococcus xylosus choline [cein (cudC), glycine betaine aldehyde [nase (cudB) genes, complete cds.] [NT:CudB]
ORF Name	NTID	AAID NT AA score probability
A17503000985_6101063_c2_1925	1291	
Description NO-HIT		
ORF Name	NTID	AAID NT AA score probability
AI7503000985_6101542_f2_836	1292	5064 132 43
Description NO-HIT		
ORF Name	NTID	AAID NT AA score probability
AI7503000985_6132893_c3_2259	1293	5065 462 153
Description NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000985_6136050_c2_1807	1294	5066 246 81 80 0.0033
[GN:TM0050] [OR:Thermotoga maritima]	[DB:ge	2:AE000512] [PN:iron(II) transport protein A] enpept-bct2] [DE:Thermotoga maritima section ar to SP:P33649 PID:414746 PID:606343]
ORF Name	NTID	AAID NT AA score probability
A17503000985_6136527_c1_1463	1295	5067 615 204 618 2.4e-60
[CL:adenylylsulfate kinase:adenylyls [DB:pir2] >gp:[GI:e332186:g2462959] 5-phosphosulfate kinase] [GN:ylnC] [[DE:Bacillus subtilis pyrE to yloA g >gp:[GI:e1185152:g2633933] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-b	sulfate [LN:BSP] OR:Baci jene reg [009] [A	Alfate kinase homolog ylnC] [GN:ylnC] kinase homology] [OR:Bacillus subtilis] PYREYLO] [AC:AJ000974] [PN:putative adenosine Allus subtilis] [DB:genpept-bct1] yion.] [LE:3535] [RE:4128] [DI:direct] AC:Z99112:AL009126] [GN:ylnC] [FN:unknown] DE:Bacillus subtilis complete genome (section Alar to adenylylsulfate kinase] [LE:34348]
ORF Name	NTID	AAID NT AA score probability
A17503000985_6251592_c2_1940	1296	5068 150 49
Description NO-HIT		

ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000985_6289068_c2_1750	1297	5069	219 72 233 1.0e-18
Description gp:[GI:g4574118] [LN:AF009415] [AC:AR [OR:Staphylococcus xylosus] [DB:genpe transporter (cudT), putativeregulator dehydrogenase(cudA), and choline dehy [LE:811] [RE:2433] [DI:direct]	ept-bct ry prot	2] [DE ein (c	E:Staphylococcus xylosus choline
ORF Name	NTID	AAID	NT AA LengthLength score probability
AI7503000985_6369688_f2_562	1298	5070	915 304 130 5.9e-06
	opsis t II BAC	haliana	PN:putative phosphatidic acid na] [SR:thale cress] [DB:genpept-pln2] genomic sequence,complete sequence.]
ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000985_6442202_f3_1033 Description	1299	5071	1050 349 310 1.0e-27
KD PROTEIN IN RPSF-SPOOJ INTERGENIC F [AC:I40448:S66018:C70084:S18084] [PN origin region)] [GN:yyaD] [OR:Bacil] [LN:BAC180K] [AC:D26185] [PN:unknown] (sub_species:Marburg, strain:168) DNA kilobase region of replication origin >gp:[GI:g580907] [LN:BSORIGS] [AC:X62 [DE:B.subtilis genes rpmH, rnpA, 50kd	REGION] N:CONSELUS SUB LUS SUB A] [DB: A] [LE 2539] [d d, gidA L:direc nknown] e (sect	[SP:P: rved hy tilis] acillus genpept :51420 OR:Bac: and g: t] >gp [OR:Bacion 21	us subtilis] [SR:Bacillus subtilis ot-bct1] [DE:B. subtilis DNA, 180 D] [RE:52436] [DI:complement] cillus subtilis] [DB:genpept-bct1] gidB.] [NT:unnamed protein product] p:[GI:e1184820:g2636641] [LN:BSUB0021] Bacillus subtilis] [DB:genpept-bct1] L of 21): from 3999281to 4214814.]
ORF Name	NTID	AAID	NT AA probability
A17503000985_651527_c1_1369	1300	5072	732 243 359 6.7e-33
[PN:hypothetical protein] [CL:N-(5'-p (5''-phosphoribosyl)-4-imidazolecarbo 6803, , PCC 6803] [SR:PCC 6803,] [DE	bhospho examide es:pir2] esilform:	DB:swis -D-ribo isomen >gp:[0 imino-5	.ssprot] >pir:[LN:S76756] [AC:S76756] oosylformimino)-5-amino-1-erase] [OR:Synechocystis sp.] [SR:PCC [GI:d1019401:g1653757] [LN:D90916] observation of the second content of th
ORF Name	NTID	AAID	NT AA LengthLength score probability
	1301	5073	138 45
Description NO-HIT			

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000985_6642792_±3_978	1302	5074		48		
Description					(
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
AI7503000985_6662875_f1_382	1303	5075	147	48		
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> <u>Length</u>	<u>AA</u> Length	score	probability
AI7503000985_6678507_f1_253	1304	5076		68		
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000985_6745327_f3_1336	1305	5077	144	47		
Description						
NO-HIT					 .	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000985_6772250_c1_1380	1306	5078		46		
Description		L				
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000985_6829635_c2_1695	1307	5079	132	43		
Description						
NO-HIT			_			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000985_6829812_c1_1542	1308	5080	948	315	795	4.2e-79
Description					············	
gp:[GI:g4959404] [LN:AF115391] [AC:A						
[OR:Lactobacillus sakei] [DB:genpept partial cds; LaaB (laaB), putative ac						
completecds; rbs operon, complete se						
[LE:4450] [RE:5358] [DI:direct]						
ORF Name	NTID	AAID	NT Length:	<u>AA</u> Length	score	probability
A17503000985_6837938_f1_376	1309	5081		1186	198	2.1e-14
Description						· · · · · · · · · · · · · · · · · · ·
sp: [LN:CDR1_HUMAN] [AC:P51861] [GN:C						
[DE:CEREBELLAR-DEGENERATION-RELATED . >pir:[LN:A29770] [AC:A29770:A35640]						-
[GN:CDR1:CDR] [OR:Homo sapiens] [SR						

NT ORF Name NTID AAID probability LengthLength A17503000985 6844012 c2 1748 1310 5082 1344 2.9e-57 Description sp:[LN:DCUA_ECOLI] [AC:P04539] [GN:DCUA:GENA] [OR:ESCHERICHIA COLI] [DE:ANAEROBIC C4-DICARBOXYLATE TRANSPORTER DCUA] [SP:P04539] [DB:swissprot] >pir:[LN:QOEC94] [AC:S56366:S57340:H65223:A04471:S08589] [PN:dicarboxylate membrane-transporter protein A:anaerobic c4-dicarboxylate transporter dcua:aspartase membrane transport protein genA] [GN:dcuA:genA] [CL:dicarboxylate membrane-transporter protein A] [OR:Escherichia coli] [DB:pirl] [MP:94 min] >gp:[GI:g510888] [LN:ECDUCA] [AC:X79887] [PN:dicarboxylate membrane-transporter protein] [GN:dcuB] [OR:Escherichia coli] [DB:genpept-bct1] [DE:E.coli dcuA gene.] [SP:P04539] [LE:129] [RE:1430] [DI:direct] >qp:[GI:q536982] [LN:ECOUW93] [AC:U14003] [GN:genA] [FN:membrane transport of aspartase] [OR:Escherichia coli] [DB:genpept-bct1] [DE:Escherichia coli K-12 chromosomal region from 92.8 to 00.1 minutes.] [LE:56300] [RE:57601] [DI:complement] >gp:[GI:g1790580] [LN:AE000486] [AC:AE000486:U00096] [PN:anaerobic dicarboxylate transport] [GN:dcuA] [FN:transport; Transport of small molecules:] [OR:Escherichia coli] [DB:genpept-bct2] [DE:Escherichia coli K-12 MG1655 section 376 of 400 of the completegenome.] [NT:f433; 100 pct identical amino acid sequence and] [LE:6845] [RE:8146] [DI:complement] NT AΑ ORF Name NTID AAID score probability LengthLength AI7503000985_6897133_f1_415 1311 5083 126 41 0.038 Description gp:[GI:g5306168] [LN:AF160864] [AC:AF160864] [PN:orf256] [GN:orf256] [OR:Mitochondrion Tetrahymena pyriformis] [SR:Tetrahymena pyriformis] [DB:genpept] [DE:Tetrahymena pyriformis mitochondrial DNA, complete genome.] [NT:Open reading frame ymf62 (CPGN)] [LE:40866] [RE:41636] [DI:direct] NT AΑ

NTID

1312

AAID

5084

score

LengthLength

280

843

probability

ORF Name

Description NO-HIT

A17503000985_6928_c1_1546

ORF Name	NTID	AAID	NT LengthLe	AA ength score	probability
AI7503000985_7039051_c3_2238	1313	5085	750 2	49 585 7	.6e-57
Description					Ti i
sp:[LN:YCKJ_BACSU] [AC:P42200] [GN:YABC TRANSPORTER PERMEASE PROTEIN] [SINC:I40451:D69761:S52382] [PN:glutaryckJ:ABC-type transport system probated [CL:histidine permease protein M] [CI:LN:BSPAAT] [AC:X77636] [PN:putative subtilis] [DB:genpept-bct1] [DE:B.suting [NT:potential ABC-transport system] >gp:[GI:e1182312:g2632646] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-bct2] [DE:Bacillus subtilis] [DB:genpept-bc2] [CI:e1182312:g2632646] [RE:215600] [LN:BSUB0003] [AC:Z99106:AL009126] [DB:genpept-bct1] [DE:Bacillus subtite to [NT:similar to glutamine] [RE:7500] [DI:complement] >gp:[GI:d1 [PN:homologue of glutamine transport [SR:Bacillus subtilis (strain:168 transport system] [DE:Bacillus subtilis (strain:168 transport system] [DI:complement] [SR:Bacillus subtilis (strain:168 transport system] [DI:complement] [SR:Bacillus subtilis (strain:168 transport system] [DI:complement]	SP:P4220 Imine AB Ible membran Ibtilis [SP:P423 IO02] [A IOC1] [D ICSIMILA IDI:COM IDI:COM	DI [DB C trans brane solutation of the color	:swisspresporter spanning btilis] nning sulve amino LE:909] 05:AL0099 llus subs lutamine t] >gp:[0 unknown] genome (ser (permedial) [LN:1 yckJ] [OI :genpept	ot] >pir:[LN (permease) Protein] [G [DB:pir2] >g bunit] [GN:C acid transport [GN:yck tilis completed ABC transport [GR:Bacillus section 3 of ease)] [SP:E [D50453] [AC:Bacillus sebtt] [DE:E [DE	J:I40451] momolog GN:yckJ] gp:[GI:g666982] DRF2] [OR:Bacillus morter gene.] DI:direct] EJ] [FN:unknown] morter (permease)] g2632661] ms subtilis] E21): from 402751 P42200] [LE:6796] D50453] subtilis] Bacillus subtilis
ORF Name	NTID	AAID	NT	AA GGOYO	probability
			LengthLe		probability
A17503000985_7227175_c3_2052	1314	5086	189 6	2	
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> LengthLe	AA score	probability
A17503000985_7228517_c3_2197	1315	5087			.3e-15
Description		L			
pir:[LN:A70556] [AC:A70556] [PN:pro [CL:mutT domain homology] [OR:Mycoba >gp:[GI:e317133:g2117198] [LN:MTCI65 [OR:Mycobacterium tuberculosis] [DB:complete genome; segment 50/162.] [N [LE:24794] [RE:25219] [DI:direct]	cterium] [AC:Z: genpept	tubero 95584: <i>1</i> -bct1]	culosis] AL123456] [DE:Myco	[DB:pir2] [PN:mutT2] obacterium t	[GN:mutT2] uberculosis H37Rv
ORF Name	NTID	AAID	<u>NT</u> LengthLe	AA ength score	probability
AI7503000985_7234627_c1_1580	1316	5088	210 63	9	
Description					
NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> LengthLe	AA ength score	probability
AI7503000985_7242128_c2_1926	1317	5089	345 13	352 3	.7e-32
Description					
<pre>gp:[GI:e314909:g2073521] [LN:SSK1MEC [OR:Staphylococcus sciuri] [DB:genpe 179.] [NT:ORF450] [LE:<1] [RE:1351]</pre>	pt-bct1]	[DE:S	.sciuri	_	

ORF Name	NTID	AAID	<u>NT</u> Length Le	AA score	probability
AI7503000985_7243832_c2_1644	1318	5090		73 684	2.4e-67
Description	L				
<pre>gp:[GI:d1044599:g5105500] [LN:AP0000 protein] [GN:APE1810] [OR:Aeropyrum [DB:genpept] [DE:Aeropyrum pernix ge site motif A (P-loop)] [LE:155809] </pre>	pernix]	[SR:A NA, se	eropyrum ction 5/	pernix (s 7.] [NT:mo	train:K1) DNA]
ORF Name	NTID	AAID	NT LengthLe	AA ength	probability
AI7503000985_7245377_c1_1534	1319	5091	1002 3	33 258	3.4e-22
Description sp:[LN:EST_ACICA] [AC:P18773] [GN:ES [DE:ESTERASE,] [SP:P18773] [DB:swiss [AC:M24890:L20754] [PN:esterase] [OR calcoaceticus ssp. lwoffii (strain R calcoaceticus esterase gene, complet	sprot] >9 R:Acinete RAG-1) D	gp:[GI obacte NA] [D	:g303953] r calcoad B:genpept	[LN:ACCE ceticus] [c-bct1] [D	STERAS] SR:Acinetobacter E:Acinetobacter
ORF Name	NTID	AAID	NT LengthLe	AA ength	probability
A17503000985_7301078_c3_2006	1320	5092		340	7.0e-31
Description			┛└ ────┛└─		
gp:[GI:g3114664] [LN:AF061267] [AC:A [OR:Pseudomonas stutzeri] [DB:genper alpha-ketoglutarate-dependenthypopho component HtxB(htxB), inner membrane HtxD(htxD), inner membrane component (htxF), HtxG (htxG), and HtxH (htxH) HtxI (htxI) gene, partial cds.] [NT:[LE:3996] [RE:5021] [DI:direct]	ot-bct2] osphite of compone HtxE (] genes,	[DE:Pdioxygent HthtxE), compl	seudomona enase (ht xC (htxC) putativa etecds; a	as stutzer EXA), bind , ATPase e C-P lyas and putati	i putative ing protein component esubunits HtxF ve C-P lyase subunit
ORF Name	NTID	AAID	<u>NT</u> LengthLe	AA ngth score	probability
A17503000985_783375_£2_893	1321	5093	147 48		
Description NO-HIT					
ORF Name	NTID	AAID	NT LengthLe	AA ngth score	probability
A17503000985_784387_c3_1970	1322	5094	1425 47		1.3e-77
Description		<u> </u>	JL		
pir:[LN:G71641] [AC:G71641] [PN:dih [GN:pdhD:RP805] [CL:dihydrolipoamid homology] [OR:Rickettsia prowazekii] [AC:AJ235273:AJ235269] [PN:DIHYDROLI [OR:Rickettsia prowazekii] [DB:genpe complete genome; segment4/4.] [LE:13	le dehydi [DB:pii POAMIDE pt-bct1]	rogena r2] >g DEHYD [DE::	se:dihydr p:[GI:e13 ROGENASE Rickettsi	colipoamid 43076:g38 (pdhD)] [.a prowaze	e dehydrogenase 61332] [LN:RPXX04] GN:RP805] kii strain Madrid E,

			NT	AA		
ORF Name	NTID	AAID	Length		score	probability
AI7503000985_787677_£2_636	1323	5095	483	160	246	6.4e-21
Description						
<pre>gp:[GI:g2735506] [LN:SCU96107] [AC:t] [OR:Staphylococcus carnosus] [DB:genus, N10-methylenetetrahydromethanopte putative transmembraneprotein genes, NhaC(nhaC) gene, partial cds.] [NT:t] [DI:direct]</pre>	npept-bo erinredu comple	t2] [D ctase te cds	E:Staph homolog , and p	ylococ , SceE outativ	cus ca precu re Na+/	rnosus rsor (sceB) and H+ antiporter
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
AI7503000985_798838_c3_1987	1324	5096	1260	419	845	2.1e-84
Description	 	<u> </u>			, (
pir:[LN:E70368] [AC:E70368] [PN:hisdehydrogenase:histidinol dehydrogenasesp:[GI:g2983343] [LN:AE000707] [AC:[GN:hisD] [OR:Aquifex aeolicus] [DB:109 of the complete genome.] [LE:571	ase homo :AE00070 :genpept	ology] 7:AE00 -bct2]	OR:Aqu] [OE:Aq [DE:Aq]	ifex a PN:his uifex	eolicu tidino aeolic	s] [DB:pir2] l dehydrogenase]
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
AI7503000985_823518_f1_169	1325	5097	225	74		
Description				-	•	
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000985_832561_c3_2102	1326	5098	138	45		
Description					•	
NO-HIT	-9-2					
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000985_86088_£2_835	1327	5099	369	122	154	3.6e-11
Description gp:[GI:g2735504] [LN:SCU96107] [AC:U [DB:genpept-bct2] [DE:Staphylococcus N5,N10-methylenetetrahydromethanopte putative transmembraneprotein genes, NhaC(nhaC) gene, partial cds.] [NT:C	carnos rinredu comple	us ctase l te cds	nomolog	, SceB utativ	precu e Na+/1	rsor (sceB) and H+ antiporter
ORF Name	NTID	AAID	<u>NT</u> Length:	<u>AA</u> Length	score	probability
AI7503000985_860917_£3_1170	1328	5100		66	56	0.0095
Description						
pir:[LN:A71605] [AC:A71605] [PN:pro [GN:PFB0845w] [OR:Plasmodium falcip [AC:AE001420:AE001362] [PN:predicted [OR:Plasmodium falciparum] [SR:malar [DE:Plasmodium falciparum chromosome [NT:predicted by GlimmerM] [LE:9688]	earum] [l integr ria para e 2, sec	DB:pir: al meml site P tion 5'	2] >gp: orane p falci 7 of 73	[GI:g3 rotein parum] of the	845291]] [GN:] [DB:ge] [LN:AE001420] PFB0845w] enpept-inv2]

ORF Name	NTID	AAID	NT AA LengthLength score	probability
AI7503000985_867255_f1_235	1329	5101	153 50	
Description				
NO-HIT				
ORF Name	NTID	AAID	NT AA LengthLength	probability
A17503000985_882143_c2_1927	1330	5102	255 84	
Description NO-HIT				
ORF Name	NTID	AAID	NT AA LengthLength score	probability
AI7503000985_891000_c2_1906	1331	5103	147 48	
Description				
NO-HIT		4.00		
ORF Name	NTID	AAID	NT AA LengthLength	probability
A17503000985_895253_c3_2019	1332	5104	510 169 303	5.8e-27
Description pir:[LN:E71960] [AC:E71960] [PN:pro				
[GN:jhp0210] [OR:Helicobacter pylor [DB:pir2] >gp:[GI:g4154731] [LN:AE00 METHIONINE SULFOXIDE REDUCTASE] [GN: [DB:genpept-bct2] [DE:Helicobacter p completegenome.] [NT:similar to H. p [DI:direct]	1459] [<i>[</i> jhp0210] ylori, s	AC:AE0 [OR: strain	01459:AE001439] [PN: Helicobacter pylori J99 section 20 of 1	putative PEPTIDE: J99] 132 of the
ORF Name	NTID	AAID	NT AA LengthLength score	probability
AI7503000985_901377_c2_1785	1333	5105		
Description				
NO-HIT				
ORF Name	NTID	AAID	NT AA LengthLength	probability
A17503000985_901377_£2_596	1334	5106	129 42	
Description				
NO-HIT				
ORF Name	NTID	AAID	NT AA LengthLength	probability
AI7503000985_953142_f1_219	1335	5107	954 317 228 5	5.1e-19
Description				
sp:[LN:Y0BS_MYCTU] [AC:Q50648] [GN:M' [DE:HYPOTHETICAL 26.2 KD PROTEIN CY2: [AC:D70724] [PN:hypothetical protein tuberculosis] [DB:pir2] >gp:[GI:e256:[PN:hypothetical protein Rv2573] [GN [DB:genpept-bct1] [DE:Mycobacterium 114/162.] [NT:Rv2573, (MTCY227.28c),	27.28C] n Rv2573 179:g147 :Rv2573] tubercul	[SP:Q 3] [GN 78239] [OR:	50648] [DB:swissprot :Rv2573] [OR:Mycoba [LN:MTCY227] [AC:Z7 Mycobacterium tubero H37Rv complete genom	c] >pir:[LN:D70724] acterium 77724:AL123456] culosis] ne; segment

ORF Name	NTID	AAID NT AA probability LengthLength score probability
A17503000985_953930_f2_838	1336	5108 132 43
Description	L	
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000985_961562_c1_1401	1337	5109 186 61
Description		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
AI7503000985_968800_c2_1796	1338	5110 804 267 1031 4.2e-104
Description		
<pre>gp:[GI:e244971:g1340128] [LN:SA1234] [DB:genpept-bct1] [DE:S.aureus orfs [DI:direct]</pre>		
ORF Name	NTID	AAID NT AA score probability
A17503000985_9766375_£1_90	1339	5111 162 53
Description NO-HIT		
ORF Name	NTID	AAID NT AA score probability
AI7503000985_9770801_f2_621	1340	5112 138 45
Description NO-HIT		
ORF Name	NTID	AAID <u>NT AA</u> score probability
A17503000985_978965_c3_2055	1341	5113 213 70
Description		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
AI7503000985_986312_f2_447	1342	5114 168 55
Description		
NO-HIT		

ORF Name	NTID	AAID	NT AA	score	probability
Description sp:[LN:DHAL_VIBCH] [AC:P23240] [GN:ADEHYDROGENASE,] [SP:P23240] [DB:swister [PN:aldehyde dehydrogenase] [GN:aldate [DB:genpept-bct1] [EC:1.2.1.3] [DE:Vomplete cds, and tagAgene, 5' end.] [LN:AF034434] [AC:AF034434] [PN:aldet [DB:genpept-bct2] [DE:Vibrio cholerate transposase, aldehyde dehydrogenase (putative inner membrane protein, and toxR-activated gene Dprotein (tagRE:3908] [DI:complement]	ALDA] [0 ssprot] A] [OR:V /ibrio o [LE:41 shyde de ae patho (aldA),	DR:VIBR >gp:[G Vibrio cholera 19] [RE chydrog ogenici toxR-a ive zin	Length Length [1533] [510] IO CHOLERAE] I:g155276] cholerae] [5] e aldehyde of :1939] [DI:of enase] [GN:a ty island, p ctivated ger cmetalloprot	[EC:1.2 [LN:VIBTA SR:V. cho dehydrogedirect] : aldA] [OF outative de A prot	2.1.3] [DE:ALDEHYDE AGALDA] [AC:M60658] clerae DNA] enase gene, egp:[GI:g3004925] R:Vibrio cholerae] cein(tagA), nes, complete cds;
ORF Name	NTID	AAID	NT AA LengthLengt	h score	probability
A17503000985_9928200_f1_189	1344	5116	159 52		
Description NO-HIT				40000	
ORF Name	NTID	AAID	NT AA LengthLengt	h score	probability
AI7503000985_9944635_f2_631	1345	5117	135 44		
Description NO-HIT					
ORF Name	NTID	AAID	NT AA LengthLengt	h score	probability
A17503000985_995300_f1_28	1346	5118	204 67	70 0	.048
Description gp:[GI:g1173895] [LN:PFU41075] [AC:Ufalciparum] [SR:malaria parasite str [DB:genpept-inv2] [DE:Plasmodium fal 1(RAP-1) gene, partial cds.] [NT:rho [LE:<1] [RE:>461] [DI:direct]	rain=Ind .ciparum	lJ-1 (c n strai	lone 1 of In n IndJ-1 rho	dian isc ptry-ass	plate J)] sociated protein
ORF Name	NTID	AAID	NT AA LengthLengt	h score	probability
AI7503000985_9954012_f1_333	1347	5119	141 46		
Description				_	

ORF Name	NTID AA	ID NT LengthI	AA Length score	probability
AI7503000986_10317307_£2_121	1348 513			0.016
Description Sp: [LN:TAGB_BACSU] [AC:P27621] [GN:TBIOSYNTHESIS PROTEIN B PRECURSOR] [S [AC:C49757:C69720] [PN:polyglycerol [GN:tagB] [OR:Bacillus subtilis] [D [AC:M57497] [GN:tagB] [OR:Bacillus subtilis] tag [DB:genpept-bct1] [DE:B.subtilis tag [NT:putative] [LE:1603] [RE:2748] [D [AC:Z99122:AL009126] [GN:tagB] [FN:p [OR:Bacillus subtilis] [DB:genpept-bl9 of 21): from 3597091to 3809700.] Sp:[GI:e1184482:g2636102] [LN:BSUB0 [FN:polyglycerol phosphate assembly [DE:Bacillus subtilis complete genom [SP:P27621] [LE:84143] [RE:85288] [D	AGB] [OR:BAP:P27621] phosphate B:pir2] >gp ubtilis] [SA, tagB, ta I:direct] >colyglycerol ct1] [DE:Ba [SP:P27621] 019] [AC:Z9 and export] e (section	ACILLUS SUR [DB:swisspicechoic acomplete acomp	BTILIS] [DE: rot] >pir:[L rid biosynth 725] [LN:BAC lis (strain gD genes, co 184482:g2636 e assembly a ptilis compl B] [RE:85288 9126] [GN:ta llus subtili	TEICHOIC ACID N:C49757] Lesis protein tagB] TAGABCD] 168) DNA] Implete cds.] 102] [LN:BSUB0019] Ind export] ete genome (section)] [DI:direct] gB] s] [DB:genpept]
ORF Name	NTID AA	ID NT LengthL	AA score	probability
A17503000986_10626525_f1_58	1349 512			1.1e-80
pir:[LN:E69831] [AC:E69831] [PN:con [CL:Bacillus subtilis conserved hypo [DB:pir2] >gp:[GI:e1183034:g2633368] [FN:unknown] [OR:Bacillus subtilis] genome (section 6 of 21): from 99950 [LE:106496] [RE:107488] [DI:direct] [PN:hypothetical protein] [GN:yhfP] [DE:B.subtilis chromosomal DNA, regi- similarity to yhdH from E.coli (Swis	thetical pr [LN:BSUB00] [DB:genpept] to1209940 pgp:[GI:e32] [OR:Bacilluon 78-80 de	rotein yhff 006] [AC:Z9 1-bct1] [DE 0.] [NT:sim 4995:g2226 as subtilis grees: apr	P] [OR:Bacil P9109:AL0091 E:Bacillus s milar to hyp 5252] [LN:BS B] [DB:genpe EE to comK.]	lus subtilis] 26] [GN:yhfP] ubtilis complete othetical proteins] Y14084] [AC:Y14084] pt-bct1] [NT:high
ORF Name	NTID AA	<u>NT</u> LengthL	AA ength score	probability
AI7503000986_10976625_c1_226	1350 512			4.8e-87
Description Sp: [LN:YHXD_BACSU] [AC:P40398:007554] [DE: (EC 1) (ORFY)] [SP:P40398:007554] [AC:E69835:S43612] [PN:ribitol dehyelochol dehydrogenase homology] [OR:Dep: [GI:e1183045:g2633379] [LN:BSUB0] [OR:Bacillus subtilis] [DB:genpept-best of 21): from 999501 to1209940.] [N' [LE:117706] [RE:118605] [DI:complementation of the complementation of the compleme	DO7554] [DE drogenase hacillus subo6] [AC:Z9ct1] [DE:Baf:similar the continuity of t	s:swissprot domolog yhx dbtilis] [E 19109:AL009 dcillus sub do ribitol d:e324973:g [OR:Bacil domal DNA,	Pjir:[LN: CD] [GN:yhxD CB:pir2] C126] [GN:yh Ctilis compl dehydrogena C2226172] [Li Lus subtili region 92 de	E69835]] [CL:short-chain xD] [FN:unknown] ete genome (section se] [SP:P40398] N:BSY14081] s] egrees:
DRF Name	NTID AAI	D NT LengthL	AA ength score	probability
A17503000986_11114677_±2_73	1351 512		57	
Description				

ORF Name	NTID	AAID	NT AA LengthLength	probability
AI7503000986_11740778_f2_80	1352	5124	216 71	
Description				
NO-HIT				
ORF Name	NTID	AAID	<u>NT AA</u> LengthLength	probability
A17503000986_1207250_c3_313 Description	1353	5125	159 89 0	.032
sp:[LN:ATP6_APIME] [AC:Q00275] [GN:A [EC:3.6.1.34] [DE:ATP SYNTHASE A CHA pir:[LN:A42622] [AC:A42622:S52964] [CL:H+-transporting ATP synthase prohoneybee] [EC:3.6.1.34] [DB:pir2] >g subunit 6] [OR:Mitochondrion Apis me [DB:genpept-inv1] [DE:Apis mellifera [LE:4584] [RE:5264] [DI:direct] >gp: [PN:adenosine triphosphatase 6] [GN:ligustica] [SR:Mitochondrion Apis me [DB:genpept-inv1] [DE:Apis mellifera (ATPase8)gene, complete cds; adenosi [LE:141] [RE:821] [DI:direct]	IN, (PRO [PN:H+- tein 6] p:[GI:g5 llifera ligusti [GI:g552 ATPase6] llifera mitocho	OTEIN 6 -transp [OR:mi 552442] ligust ca com 2451] [OR:M	[DB: Q00275] [DB: corting ATP synthase tochondrion Apis me [LN:AMFGENOM] [AC: cica] [SR:common home plete mitochondrial [LN:AMFMTATPAS] [AC: Mitochondrion Apis me cica (organelle Mitochosine triphosp	swissprot] e, chain 6] ellifera] [SR:, L06178] [PN:ATPase eybee] egenome.] M87065] ellifera chondrio] chatase 8
ORF Name	NTID		NT AA LengthLength	probability
A17503000986_1290703_c1_227 Description NO-HIT	1354	5126	[141] [46]	
ORF Name	NTID	AAID	NT <u>AA</u> LengthLength	probability
A17503000986_14644018_f1_39	1355	5127	969 322 324 3	.4e-29
Description gp:[GI:g2196513] [LN:SEU77778] [AC:U [GN:epiH] [FN:involved in epidermin [DB:genpept-bct1] [DE:Staphylococcus transportersubunits (epiG), (epiE), (epiT') and EpiT'' (epiT'') genes, c [DI:direct]	secretic epiderm and (epi	on] [OR nidis p .F), pu	plasmid pTue32 putat stative membrane pro	dermidis] ive ABC tein(epiH), EpiT'
ORF Name	NTID	<u>AAID</u>	NT AA LengthLength	probability
A17503000986_14647750_c2_238	1356	5128	141 46	
Description				
NO-HIT				
ORF Name A17503000986_14656952_t2_93 Description	NTID 1357	<u>AAID</u> 5129	NT AA LengthLength 633 210	probability

ORF Name	$rac{ ext{NTID}}{ ext{Length}} rac{ ext{AA}}{ ext{Length}} rac{ ext{Score}}{ ext{probability}}$
AI7503000986_15035952_£2_91	1358 5130 477 158
Description NO-HIT	
ORF Name	NTID AAID NT AA score probability
AI7503000986_15117317_c2_243	1359 5131 2184 727 953 2.4e-102
ACID BIOSYNTHESIS PROTEIN F] [SP:P13 [AC:S06049:G69720] [PN:probable CDP:CDP-glycerol:polyglycerol phosphate protein:teichoic-acid synthase] [GN: [EC:2.7.8.12] [DB:pir2] [MP:310 degr [OR:Bacillus subtilis] [DB:genpept-b (tag3) polypeptide (AA 1-746)] [SP:P>gp:[GI:e1184478:g2636098] [LN:BSUB0 [PN:CDP-glycerol:polyglycerol phosph [OR:Bacillus subtilis] [DB:genpept-b 19 of 21): from 3597091to 3809700.] [LE:78129] [RE:80369] [DI:complement [AC:Z99122:AL009126] [PN:CDP-glycero acid biosynthesis] [OR:Bacillus subt	e glycero-phosphotransferase tagF:rodC rodC:tag-3:tagF] [OR:Bacillus subtilis] rees] >gp:[GI:g40100] [LN:BSRODC] [AC:X15200] rot1] [DE:Bacillus subtilis rodC operon.] [NT:rodC r13485] [LE:2178] [RE:4418] [DI:direct] rot1] [AC:Z99122:AL009126] rate] [GN:tagF] [FN:teichoic acid biosynthesis] rot1] [DE:Bacillus subtilis complete genome (section [NT:alternate gene name: rodC] [SP:P13485] rect] >gp:[GI:e1184478:g2636098] [LN:BSUB0019] rol:polyglycerol phosphate] [GN:tagF] [FN:teichoic cilis] [DB:genpept] [DE:Bacillus subtilis complete rog01 3809700.] [NT:alternate gene name: rodC]
ORF Name	$ ext{NTID}$ $ ext{AAID}$ $ ext{LengthLength}$ $ ext{score}$ $ ext{probability}$
A17503000986_15136562_c2_253	1360
[SR:Bacillus halodurans (strain:C-12	[AC:AB013369] [OR:Bacillus halodurans] [DB:genpept-bct1] [DE:Bacillus halodurans andcomplete cds.] [NT:unknown] [LE:4328] [RE:5830]
gp:[GI:d1039113:g4514332] [LN:AB0133 [SR:Bacillus halodurans (strain:C-12 C-125 yesT and comEC genes, partial	25) DNA] [DB:genpept-bct1] [DE:Bacillus halodurans
<pre>gp:[GI:d1039113:g4514332] [LN:AB0133 [SR:Bacillus halodurans (strain:C-12 C-125 yesT and comEC genes, partial [DI:direct] ORF Name A17503000986_15829135_c2_237</pre>	25) DNA] [DB:genpept-bct1] [DE:Bacillus halodurans andcomplete cds.] [NT:unknown] [LE:4328] [RE:5830] NTID AAID NT AA Score probability
gp:[GI:d1039113:g4514332] [LN:AB0133 [SR:Bacillus halodurans (strain:C-12 C-125 yesT and comEC genes, partial [DI:direct] ORF Name A17503000986_15829135_c2_237 Description pir:[LN:E69764] [AC:E69764] [PN:hyp subtilis] [DB:pir2] >gp:[GI:e1182361 [GN:ycnI] [FN:unknown] [OR:Bacillus complete genome (section 3 of 21): f [DI:complement] >gp:[GI:d1009660:g18 [OR:Bacillus subtilis] [SR:Bacillus	NTID AAID NT AA LengthLength Score probability NTID AAID LengthLength Score probability December
gp:[GI:d1039113:g4514332] [LN:AB0133 [SR:Bacillus halodurans (strain:C-12 C-125 yesT and comEC genes, partial [DI:direct] ORF Name A17503000986_15829135_c2_237 Description pir:[LN:E69764] [AC:E69764] [PN:hyp subtilis] [DB:pir2] >gp:[GI:e1182361 [GN:ycnI] [FN:unknown] [OR:Bacillus complete genome (section 3 of 21): f [DI:complement] >gp:[GI:d1009660:g18 [OR:Bacillus subtilis] [SR:Bacillus [DE:Bacillus subtilis] DNA for 25-36	NTID AAID NT AA LengthLength Score probability NTID AAID LengthLength Score probability December
<pre>gp:[GI:d1039113:g4514332] [LN:AB0133 [SR:Bacillus halodurans (strain:C-12 C-125 yesT and comEC genes, partial [DI:direct] ORF Name A17503000986_15829135_c2_237 Description pir:[LN:E69764] [AC:E69764] [PN:hyp subtilis] [DB:pir2] >gp:[GI:e1182361 [GN:ycnI] [FN:unknown] [OR:Bacillus complete genome (section 3 of 21): f [DI:complement] >gp:[GI:d1009660:g18 [OR:Bacillus subtilis] [SR:Bacillus [DE:Bacillus subtilis] DNA for 25-36 complete cds.] [LE:125407] [RE:12602</pre>	NTID AND [DB:genpept-bct1] [DE:Bacillus halodurans andcomplete cds.] [NT:unknown] [LE:4328] [RE:5830] NTID AAID NT AA Score probability 1361 5133 204 67 105 1.0e-05

ORF Name AI7503000986_17047575_f3_150 Description	NTID 1363	<u>AAID</u>	NT AA LengthLength 171 56
NO-HIT ORF Name	NTID	AAID	NT AA score probability
A17503000986 174218 cl 209	1364	5136	Length Length 1225 1.1e-124 1.25 1.1e-124 1.25 1.1e-124 1.25 1.1e-124 1.25 1.1e-124 1.25 1
Description			
sp:[LN:YHCA_BACSU] [AC:P54585] [GN:YI KD PROTEIN IN GLPD-CSPB INTERGENIC RI [AC:E69821] [PN:multidrug resistance [CL:lincomycin-resistance protein lm: >gp:[GI:e1182890:g2633224] [LN:BSUB06	EGION] e protei rB] [OR: 005] [AC ctl] [DE T:simila	[SP:P5 in hom :Bacil C:Z991 E:Baci ar to	4585] [DB:swissprot] >pir:[LN:E69821] olog yhcA] [GN:yhcA] lus subtilis] [DB:pir2] 08:AL009126] [GN:yhcA] [FN:unknown] llus subtilis complete genome (section
ORF Name	NTID	AAID	NT AA LengthLength
AI7503000986_179653_c3_325	1365	5137	
Description		L	JL
[OR:Plasmid pNS1] [SR:Plasmid pNS1 fr [DB:genpept-bct1] [DE:Plasmid pNS1 (fr	swisspro [GN:tet >gp:[GI rom Stap from Stap from Sta), compl 449] [PN 181] [DE s, pT181 [LE:1] ycline r ylococcu ne resis GI:gN:tet H1] [DB:	ot] >p] [CL]:g476 phylocophy	ir:[LN:S42238] [AC:S42238] :tetracycline resistance protein] 735] [LN:PNS1CG] [AC:M16217] [GN:tet] occus aureus, plasmid pTP5 DNA] coccus aureus) enome.] [LE:305] [RE:1684] [DI:direct] K)] [GN:tet(K)] [OR:Staphylococcus ept-bct1] [DE:tet(K)=tetracycline mid, 1380 nt].] [NT:tetracycline 380] [DI:direct] >gp:[GI:g1052998] ance protein] [GN:tet] eus plasmid pKH6] [DB:genpept-bct1] plasmid pKH6, complete sequence.] [LN:SAU38656] [AC:U38656] :Staphylococcus aureus
ORF Name AI7503000986_19548755_t3_132 Description NO-HIT	NTID 1366	<u>AAID</u> 5138	NT AA LengthLength 168 55
ORF Name A17503000986_1970178_f3_146 Description	NTID 1367	<u>AAID</u> 5139	NT AA LengthLength score probability [219] [72]

	NTID	AAID LengthLength score probability
A17503000986_19742962_c1_221	1368	5140 1026 341 423 1.1e-39
_	occus la	C:AJ011653] [PN:aldose 1-epimerase] actis] [DB:genpept-bct1] [EC:5.1.3.3] ctose operon (galAMKTEgenes).] [LE:1990]
[RE:3009] [DI:direct]		
ORF Name	NTID	AAID NT AA score probability LengthLength
AI7503000986_19960885_c1_228	1369	5141 156 51
Description NO-HIT		
ORF Name	NTID	AAID LengthLength score probability
A17503000986_20579752_c3_319	1370	5142 150 49
Description NO-HIT		
ORF Name	NTID	AAID NT AA score probability
AI7503000986_20580443_c3_310	1371	5143 333 110 173 3.5e-13
Description		
plasmid pMRC01, complete plasmidseque percent] [LE:44384] [RE:44728] [DI:d] [DB:ge ence.] [irect]	enpept-bct2] [DE:Lactococcus lactis DPC3147 [NT:similar to GB:X69895 SP:P39044 PID:40067
ORF Name	NTID	AAID LengthLength score probability
A17503000986_20718790_f3_138	1372	5144 150 49
5 ' ' '		
Description NO-HIT		
	NTID	AAID NT AA score probability
NO-HIT	<u> </u>	AAID — score probability
NO-HIT ORF Name A17503000986_20789507_c2_280 Description		Length Length probability
NO-HIT ORF Name Al7503000986_20789507_c2_280		Length Length probability
NO-HIT ORF Name A17503000986_20789507_c2_280 Description	1373	Length Length probability
NO-HIT ORF Name A17503000986_20789507_c2_280 Description NO-HIT ORF Name A17503000986_20897125_c1_212	1373 NTID	LengthLength score probability 5145 186 61
NO-HIT ORF Name A17503000986_20789507_c2_280 Description NO-HIT ORF Name	NTID 1374	AAID Length Length Score Probability 5145 186 61 AAID NT AA Score Probability

ORF Name	$\underline{\mathtt{NTID}}$	AAID LengthLength score probability
A17503000986_21676937_c1_190	1375	5147 132 43
Description		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000986_22074200_f3_184	1376	5148 126 42
Description NO-HIT		
ORF Name	NTID	AAID LengthLength score probability
A17503000986_22277215_f3_142	1377	5149 1164 387 109 1.5e-05
Description pir:[LN:C44863] [AC:C44863] [PN:R45	antige	n] [OR:Plasmodium falciparum] [DB:pir3]
ORF Name	NTID	AAID NT AA score probability
A17503000986_22455213_f3_169	1378	5150 687 228 411 2.1e-38
Description		
[GN:TM1655] [OR:Thermotoga maritima]	[DB:gen	:AE000512] [PN:response regulator DrrA] hpept-bct2] [DE:Thermotoga maritima section ilar to PID:1575577 GB:AE000512 percent]
ORF Name	NTID	AAID NT AA score probability LengthLength
		Lengthbength
A17503000986_22867942_c1_224	1379	5151 711 236 417 4.8e-39
Description gp:[GI:g4262236] [LN:ATAC006200] [ACisomerase] [GN:F10A8.17] [OR:Arabido	:AC00620 opsis that II BAC 1	5151 711 236 417 4.8e-39
Description gp:[GI:g4262236] [LN:ATAC006200] [ACisomerase] [GN:F10A8.17] [OR:Arabido [DE:Arabidopsis thaliana chromosome	:AC00620 opsis that II BAC 1	[PN:putative ribose 5-phosphate aliana] [SR:thale cress] [DB:genpept-pln2] [F10A8 genomic sequence, complete sequence.]
Description gp:[GI:g4262236] [LN:ATAC006200] [ACisomerase] [GN:F10A8.17] [OR:Arabido [DE:Arabidopsis thaliana chromosome [LE:58788] [RE:59585] [DI:complement	::AC00620 ppsis that II BAC 1	5151 711 236 417 4.8e-39 00] [PN:putative ribose 5-phosphate aliana] [SR:thale cress] [DB:genpept-pln2] F10A8 genomic sequence, complete sequence.]
Description gp:[GI:g4262236] [LN:ATAC006200] [ACisomerase] [GN:F10A8.17] [OR:Arabido [DE:Arabidopsis thaliana chromosome [LE:58788] [RE:59585] [DI:complement ORF Name AT7503000986_22869687_c1_186 Description pir:[LN:A64946] [AC:A64946] [PN:hyp protein pcoC] [OR:Escherichia coli] [AC:AE000278:U00096] [PN:orf, hypoth [OR:Escherichia coli] [DB:genpept-bc]	Price of the control	[PN:putative ribose 5-phosphate aliana] [SR:thale cress] [DB:genpept-pln2] [F10A8 genomic sequence, complete sequence.] AAID NT AA LengthLength score probability [S152] [438] [145] [143] [5.2e-10] Al protein b1841] [CL:copper resistance
Description gp:[GI:g4262236] [LN:ATAC006200] [AC isomerase] [GN:F10A8.17] [OR:Arabido [DE:Arabidopsis thaliana chromosome [LE:58788] [RE:59585] [DI:complement ORF Name AI7503000986_22869687_c1_186 Description pir:[LN:A64946] [AC:A64946] [PN:hyp protein pcoC] [OR:Escherichia coli] [AC:AE000278:U00096] [PN:orf, hypoth [OR:Escherichia coli] [DB:genpept-bc 400 of the completegenome.] [NT:f124	Price of the control	[PN:putative ribose 5-phosphate aliana] [SR:thale cress] [DB:genpept-pln2] [F10A8 genomic sequence, complete sequence.] AAID NT AA LengthLength score probability [S152] [A38] [A5] [A3] [5.2e-10 [A1] [A1] [CL:copper resistance all protein b1841] [CL:copper resistance all protein] [GN:b1841] [FN:orf; Unknown] [Escherichia coli K-12 MG1655 section 168 of 124 aa ORF is 39 pct identical (6 gaps)]
Description gp:[GI:g4262236] [LN:ATAC006200] [AC isomerase] [GN:F10A8.17] [OR:Arabido [DE:Arabidopsis thaliana chromosome [LE:58788] [RE:59585] [DI:complement ORF Name AT7503000986_22869687_c1_186 Description pir:[LN:A64946] [AC:A64946] [PN:hyp protein pcoC] [OR:Escherichia coli] [AC:AE000278:U00096] [PN:orf, hypoth [OR:Escherichia coli] [DB:genpept-bc 400 of the completegenome.] [NT:f124 [LE:4445] [RE:4819] [DI:complement]	Price of the control	[PN:putative ribose 5-phosphate aliana] [SR:thale cress] [DB:genpept-pln2] [F10A8 genomic sequence, complete sequence.] AAID NT AA LengthLength score probability [S152] [438] [145] [143] [5.2e-10] [Al protein b1841] [CL:copper resistance all protein] [GN:b1841] [FN:orf; Unknown] [Escherichia coli K-12 MG1655 section 168 of 124 aa ORF is 39 pct identical (6 gaps)]
Description gp:[GI:g4262236] [LN:ATAC006200] [AC isomerase] [GN:F10A8.17] [OR:Arabido [DE:Arabidopsis thaliana chromosome [LE:58788] [RE:59585] [DI:complement] ORF Name AT7503000986_22869687_C1_186 Description pir:[LN:A64946] [AC:A64946] [PN:hypprotein pcoC] [OR:Escherichia coli] [AC:AE000278:U00096] [PN:orf, hypoth [OR:Escherichia coli] [DB:genpept-bc400 of the completegenome.] [NT:f124 [LE:4445] [RE:4819] [DI:complement] ORF Name	Price of the control	[PN:putative ribose 5-phosphate aliana] [SR:thale cress] [DB:genpept-pln2] [Floas genomic sequence, complete sequence.] [AAID NT AA LengthLength Score probability [S152 438 145 143 5.2e-10 [AI protein b1841] [CL:copper resistance classes] [SP:[GI:g1788146] [LN:AE000278] [SP:[GI:g1788146] [FN:orf; Unknown] [Escherichia coli K-12 MG1655 section 168 of 124 aa ORF is 39 pct identical (6 gaps)] [AAID NT AA LengthLength Score probability [AAID NT AA LengthLength] [AAID LengthLength] [AAID NT AA LengthLength] [AAID LengthLength] [AAID NT AA LengthLength] [AAID NT AA LengthLength] [AAID NT AA LengthLength] [AAID LengthLength] [AAID NT AA LengthLength] [AAID LengthLength] [A

ORF Name	NTID	AAID LengthLength score probability
A17503000986_23601557_f1_10	1382	5154 201 66
Description		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000986_23601577_f3_144	1383	5155 [186] 61
Description		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000986_23611437_c1_188	1384	5156 135 44
Description		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000986_23839193_c1_193	1385	5157 1614 537 1324 3.7e-135
Description		
permease] [OR:Bacillus subtilis] [DE [AC:Z99105:AL009126] [PN:L-lactate p	B:pir2] permease lis com	plete genome (section 2 of 21): from 194651
ORF Name	NTID	AAID Length Length score probability
	NTID 1386	AAID NT AA score probability [5158 123 40
ORF Name A17503000986_23876887_f1_60 Description		LengthLength score probability
A17503000986_23876887_f1_60		LengthLength score probability
A17503000986_23876887_f1_60 Description		LengthLength score probability 5158 123 40
Description NO-HIT ORF Name	NTID	AAID LengthLength score probability 5158 123 40 AAID NT AA LengthLength score probability
Description NO-HIT ORF Name A17503000986_23923412_f1_31	1386	LengthLength score probability 5158 123 40
Description NO-HIT ORF Name	NTID	AAID LengthLength score probability 5158 123 40 AAID NT AA LengthLength score probability
Description NO-HIT ORF Name A17503000986_23923412_f1_31 Description	NTID	LengthLength score probability Sissippoint
Description NO-HIT ORF Name A17503000986_23923412_f1_31 Description NO-HIT	NTID 1387	AAID LengthLength score probability AAID NT AA Score probability AAID NT AA Score probability AAID NT AA Score probability
A17503000986_23876887_f1_60 Description NO-HIT ORF Name A17503000986_23923412_f1_31 Description NO-HIT ORF Name	NTID 1387 NTID	AAID LengthLength score probability AAID NT AA LengthLength score probability 5159 225 74 AAID NT AA LengthLength score probability LengthLength score probability
A17503000986_23876887_f1_60 Description NO-HIT ORF Name A17503000986_23923412_f1_31 Description NO-HIT ORF Name A17503000986_23929627_f3_160 Description pir:[LN:I40601] [AC:I40601:S70950]	NTID 1387 NTID 1388 [PN:mob:	AAID NT AA LengthLength Score probability AAID NT AA LengthLength 5159 225 74 AAID NT AA LengthLength EngthLength Score probability AAID LengthLength Score probability 5160 177 58 46 0.046
A17503000986_23876887_f1_60 Description NO-HIT ORF Name A17503000986_23923412_f1_31 Description NO-HIT ORF Name A17503000986_23929627_f3_160 Description pir:[LN:I40601] [AC:I40601:S70950] [OR:Bacteroides vulgatus] [DB:pir2] [PN:mobilization protein] [GN:mobA]	NTID 1388 NTID 1388 [PN:mob: >gp:[GI [FN:con:	AAID LengthLength AAID NT AA LengthLength Store probability AAID NT AA LengthLength AAID LengthLength EngthLength AAID NT AA LengthLength AAID LengthLength GN:mobA] 123 40 AAID NT AA LengthLength [GN:mobA] 124 125 127 128 128 128 128 128 128 128 128 128 128
Description NO-HIT ORF Name A17503000986_23923412_f1_31 Description NO-HIT ORF Name A17503000986_23923412_f1_31 Description NO-HIT ORF Name A17503000986_23929627_f3_160 Description pir: [LN:I40601] [AC:I40601:S70950] [OR:Bacteroides vulgatus] [DB:pir2] [PN:mobilization protein] [GN:mobA] vulgatus] [DB:genpept-bct1] [DE:Bact	NTID 1388 NTID 1388 [PN:mob: >gp:[GI [FN:con: eroides	AAID LengthLength AAID NT AA LengthLength Store probability AAID NT AA LengthLength AAID LengthLength EngthLength AAID NT AA LengthLength AAID LengthLength GN:mobA] 123 140 AAID NT AA LengthLength [GN:mobA] 124 125 127 [GN:mobA] 125 127 128 128 128 128 128 128 128 128 128 128
Description NO-HIT ORF Name A17503000986_23923412_f1_31 Description NO-HIT ORF Name A17503000986_23923412_f1_31 Description NO-HIT ORF Name A17503000986_23929627_f3_160 Description pir: [LN:I40601] [AC:I40601:S70950] [OR:Bacteroides vulgatus] [DB:pir2] [PN:mobilization protein] [GN:mobA] vulgatus] [DB:genpept-bct1] [DE:Bact	NTID 1388 NTID 1388 [PN:mob: >gp:[GI [FN:con: eroides	AAID LengthLength AAID NT AA LengthLength Store probability AAID NT AA LengthLength AAID LengthLength EngthLength AAID NT AA LengthLength AAID LengthLength GN:mobA] 123 40 AAID NT AA LengthLength [GN:mobA] 124 125 127 128 128 128 128 128 128 128 128 128 128
Description NO-HIT ORF Name A17503000986_23923412_f1_31 Description NO-HIT ORF Name A17503000986_23929627_f3_160 Description pir:[LN:I40601] [AC:I40601:S70950] [OR:Bacteroides vulgatus] [DB:pir2] [PN:mobilization protein] [GN:mobA] vulgatus] [DB:genpept-bct1] [DE:Bactcomplete cds andmobilization protein [DI:complement]	NTID 1388 NTID 1388 [PN:mob: >gp:[GI [FN:con: eroides (mobA)	AAID LengthLength Score probability AAID NT AA LengthLength Score probability AAID LengthLength Score probability AAID LengthLength Score probability AAID LengthLength 60.046 ilization protein] [GN:mobA] :g1079659] [LN:BVU38243] [AC:U38243:M72418] jugal transfer of Tn4555] [OR:Bacteroides vulgatus beta-lactamase (cfxA) gene, gene, complete cds.] [LE:1208] [RE:2611]
Description NO-HIT ORF Name A17503000986_23923412_f1_31 Description NO-HIT ORF Name A17503000986_23929627_f3_160 Description pir: [LN:I40601] [AC:I40601:S70950] [OR:Bacteroides vulgatus] [DB:pir2] [PN:mobilization protein] [GN:mobA] vulgatus] [DB:genpept-bct1] [DE:Bactcomplete cds andmobilization protein [DI:complement] ORF Name	NTID 1388 NTID 1388 [PN:mob: >gp:[GI [FN:con: eroides	AAID LengthLength Score probability AAID NT AA LengthLength Score probability AAID LengthLength Score probability AAID LengthLength Score probability AAID LengthLength Score probability [GN:mobA] [GN:mobA] [G1079659] [LN:BVU38243] [AC:U38243:M72418] [Jugal transfer of Tn4555] [OR:Bacteroides vulgatus beta-lactamase (cfxA) gene, gene, complete cds.] [LE:1208] [RE:2611] AAID NT AA LengthLength Score probability
Description NO-HIT ORF Name AT7503000986_23923412_f1_31 Description NO-HIT ORF Name AT7503000986_23929627_f3_160 Description pir:[LN:I40601] [AC:I40601:S70950] [OR:Bacteroides vulgatus] [DB:pir2] [PN:mobilization protein] [GN:mobA] vulgatus] [DB:genpept-bct1] [DE:Bact complete cds andmobilization protein [DI:complement] ORF Name AT7503000986_24250317_c1_203	NTID 1388 NTID 1388 [PN:mob: >gp:[GI [FN:con: eroides (mobA)	AAID LengthLength Score probability [GN:mobA] [GN:mobA] [GN:mobA] [G1079659] [LN:BVU38243] [AC:U38243:M72418] [Jugal transfer of Tn4555] [OR:Bacteroides vulgatus beta-lactamase (cfxA) gene, gene, complete cds.] [LE:1208] [RE:2611]
Description NO-HIT ORF Name AI7503000986_23923412_f1_31 Description NO-HIT ORF Name AI7503000986_23929627_f3_160 Description pir: [LN:I40601] [AC:I40601:S70950] [OR:Bacteroides vulgatus] [DB:pir2] [PN:mobilization protein] [GN:mobA] vulgatus] [DB:genpept-bct1] [DE:Bactcomplete cds andmobilization protein [DI:complement] ORF Name	NTID 1388 NTID 1388 [PN:mob: >gp:[GI [FN:con: eroides (mobA)	AAID LengthLength Score probability AAID NT AA LengthLength Score probability AAID LengthLength Score probability AAID LengthLength Score probability AAID LengthLength Score probability [GN:mobA] [GN:mobA] [GN:mobA] [G1079659] [LN:BVU38243] [AC:U38243:M72418] [Jugal transfer of Tn4555] [OR:Bacteroides vulgatus beta-lactamase (cfxA) gene, gene, complete cds.] [LE:1208] [RE:2611] AAID NT AA LengthLength Score probability

ORF Name	NTID	AAID	NT AA LengthLength
A17503000986_24267567_t3_163	1390	5162	645 214 99 0.0046
Description pir: [LN:C70649] [AC:C70649] [PN:hyp [OR:Mycobacterium tuberculosis] [DB: [AC:Z83866:AL123456] [PN:hypothetica tuberculosis] [DB:genpept-bct1] [DE: segment 133/162.] [NT:Rv3058c, (MTCY [DI:complement]	pir2] >q l prote: Mycobac	gp:[GI: in Rv30 terium	:e290931:g1781155] [LN:MTCY22D7] 058c] [GN:Rv3058c] [OR:Mycobacterium
ORF Name	NTID	AAID	NT AA score probability
AI7503000986_24394175_f3_151	1391	5163	159 52
Description		L	/ l
NO-HIT			
ORF Name	NTID	AAID	NT AA LengthLength
A17503000986_24407827_c2_266	1392	5164	561 186
Description	L		
NO-HIT			
ORF Name	NTID	AAID	NT AA LengthLength
A17503000986_24412826_c3_323	1393	5165	963 320 286 3.7e-25
Description			
sp:[LN:HUTG_KLEAE] [AC:P19452] [GN:H [DE:(HISTIDINE UTILIZATION PROTEIN G >gp:[GI:g149204] [LN:KPNHUTC] [AC:M3 aerogenes (strain W70) DNA] [DB:genp repressor C (hutC) gene, completecds [RE:669] [DI:direct]) (FRAGI 4604] [(ept-bct:	MENT)] DR:Kleb 1] [DE:	[SP:P19452] [DB:swissprot] psiella aerogenes] [SR:Klebsiella K.aerogenes histidine utilization
ORF Name	NTID	AAID	NT AA LengthLength
AI7503000986_24415925_c1_211	1394	5166	903 300 282 9.7e-25
Description			
<pre>gp:[GI:g1209223] [LN:ACCEST] [AC:L38 lwoffii] [DB:genpept-bct1] [DE:Acine complete cds.] [LE:638] [RE:1549] [D</pre>	tobactei	lwoff	
ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000986_24662825_c2_279	1395	5167	618 205 538 7.3e-52
Description			
pir:[LN:H70068] [AC:H70068] [PN:hypesubtilis] [DB:pir2] >gp:[GI:e1184514] [GN:ywrF] [FN:unknown] [OR:Bacillus complete genome (section 19 of 21): [DI:direct] >gp:[GI:e311284:g1929333] subtilis] [DB:genpept-bct1] [DE:B.sulywrO gene.] [LE:4588] [RE:5205] [DI:GLN:BSUB0019] [AC:Z99122:AL009126] [OB:genpept] [DE:Bacillus subtilis complete genome [LE:119966] [RE:120583] [DE:3809700.]	g263613 subtilis from 359 [LN:BS btilis I compleme GN:ywrF]	33] [LN: 57091to 5293767 DNA; 15 ent] >g [FN:u genome	genpept-bctl] [DE:Bacillus subtilis 3809700.] [LE:119966] [RE:120583] [AC:Z93767] [GN:ywrF] [OR:Bacillus 2.2 kb fragment, from ywqN gene to p:[GI:e1184514:g2636133] nknown] [OR:Bacillus subtilis]

ORF Name	NTID AAID NT AA score probability
A17503000986_2470010_c1_191	1396 5168 723 240 157 1.3e-08
falciparum] [SR:malaria parasite P.	3P5] [AC:AL034556] [GN:MAL3P5.8] [OR:Plasmodium falciparum] [DB:genpept-inv1] [DE:Plasmodium e.] [NT:predicted using hexExon; MAL3P5.8 [DI:complement]
ORF Name	NTID AAID NT AA score probability
A17503000986_24803386_c1_201	1397 5169 1080 359 215 1.4e-15
[OR:Mycobacterium tuberculosis] [DB [AC:Z77250:AL123456] [PN:hypothetic tuberculosis] [DB:genpept-bct1] [DE	pothetical protein Rv2563] [GN:Rv2563] :pir2] >gp:[GI:e1299946:g3261618] [LN:MTCY9C4] al protein Rv2563] [GN:Rv2563] [OR:Mycobacterium :Mycobacterium tuberculosis H37Rv complete genome; 9C4.05c), len: 349. Unknown membrane] [LE:5757]
ORF Name	NTID AAID LengthLength score probability
AI7503000986_2540907_c3_304	1398 5170 186 61
Description	
NO-HIT	
ORF Name	NTID AAID NT AA score probability
AI7503000986_25527188_c1_225	1399 5171 1137 378 830 8.3e-83
hydrolase] [OR:Bacillus subtilis] [IAC:Z99118:AL009126] [PN:hippurate IDB:genpept-bct1] [EC:3.5.1.32] [DE 21): from 2795131to 3013540.] [LE:20.5gp:[GI:el185802:g2635413] [LN:BSUBG [GN:hipO] [OR:Bacillus subtilis] [DI complete genome (section 16 of 21): [DI:complement] >gp:[GI:g2293256] [DI:complement] >gp:[GI:g2293256]	0016] [AC:Z99119:AL009126] [PN:hippurate hydrolase] B:genpept-bct1] [EC:3.5.1.32] [DE:Bacillus subtilis from 2997771to 3213410.] [LE:1020] [RE:2270] LN:AF008220] [AC:AF008220] [PN:putative hippurate ubtilis] [DB:genpept-bct2] [DE:Bacillus subtilis
ORF Name	NTID AAID NT AA score probability
A17503000986_25551640_c1_214	1400 5172 861 286 112 0.0023
Description	
<pre>falciparum] [SR:malaria parasite P.</pre>	3P6] [AC:Z98551] [GN:MAL3P6.11] [OR:Plasmodium falciparum] [DB:genpept-inv1] [DE:Plasmodium e.] [NT:predicted using hexExon; MAL3P6.11 [DI:complement]

ORF Name	NTID	AAID LengthLength score probability
A17503000986_25579390_f1_50	1401	5173 1194 397 402 8.5e-37
Description		
<pre>gp:[GI:e1294490:g3169038] [LN:SC1C3] [GN:SC1C3.12] [OR:Streptomyces coeli coelicolor cosmid 1C3.] [NT:SC1C3.12 [LE:13186] [RE:15279] [DI:direct]</pre>	color]	
ORF Name	NTID	AAID <u>NT AA</u> score probability
A17503000986_25910952_c3_309	1402	5174 141 46
Description NO-HIT		
ORF Name	NTID	AAID <u>NT AA</u> score probability
AI7503000986_26753588_f1_44 Description	1403	5175 1383 460 344 2.6e-31
[GN:yclK] [OR:Bacillus subtilis] [D [AC:Z99106:AL009126] [GN:yclK] [FN:u [DE:Bacillus subtilis complete genom [NT:similar to two-component sensor >gp:[GI:d1009643:g1805446] [LN:D5045 phosphatase synthesis] [GN:yclK] [OR	B:pir2] nknown] e (sect: histidir 3] [AC:I :Bacillu bct1] [I	ne kinase] [LE:24077] [RE:25498] [DI:direct] D50453] [PN:homologue of alkaline us subtilis] [SR:Bacillus subtilis DE:Bacillus subtilis DNA for 25-36 degree
ORF Name	NTID	AAID NT AA score probability
A17503000986_272550_c2_268	1404	5176 123 40
Description NO-HIT		
ORF Name	NTID	AAID LengthLength score probability
A17503000986_292883_c2_229	1405	5177 126 41
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000986_29565627_c3_286	1406	5178 879 292
Description NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000986_30476575_c1_216	1407	5179 165 54
Description NO-HIT		

ORF Name AI7503000986_30703458_c2_230 Description sp:[LN:PTSB_STAXY] [AC:P51184] [GN:S [DE:(EC 2.7.1.69) (EII-SCR)] [SP:P51 [PN:scrA protein] [CL:phosphotransfe [OR:Staphylococcus xylosus] [DB:pir2 [PN:EIIscr] [GN:scrA] [OR:Staphylococcus and unidentified open reading f [DI:direct]	184] [Derase sy 2] >gp:[occus xy	5180 90 R:STAPHYLO B:swisspro stem sucro GI:g407908 Losus] [DI	DCOCCUS X bt] >pir: pse-speci 3] [LN:SX 3:genpept	YLOSUS [LN:S3 fic en SCRA]	2.3e-126 [EC:2.7.1.69] 9978] [AC:S39978] zyme II, factor II] [AC:X69800] [DE:S.xylosus scrA
ORF_Name AI7503000986_34032561_c3_316 Description NO-HIT	NTID 1409	$\Delta \Delta TD -$	T AA gthLengtl 3 40	score	probability
ORF Name A17503000986_34199202_c1_189 Description gp:[GI:e1184294:g2635712] [LN:BSUB00 regulator] [GN:paiA] [FN:negative resubtilis] [DB:genpept-bct1] [DE:Baci from 3197001to 3414420.] [LE:106800]	gulatio .llus su	5182 54 E299120:AI of sport	L009126] plation,	343 [PN:traseptat:	3.3e-31 anscriptional ion] [OR:Bacillus
ORF Name A17503000986_34610667_c3_297 Description pir:[LN:B69377] [AC:B69377] [PN:ABC [CL:unassigned ATP-binding cassette [OR:Archaeoglobus fulgidus] [DB:pir2 [AC:AE001033:AE000782] [PN:ABC trans [OR:Archaeoglobus fulgidus] [DB:genp 172 of the complete genome.] [NT:sim [LE:2126] [RE:2812] [DI:complement]	protein } >gp:[6 porter, ept-bct	orter, ATE:: ATP-bir GI:g264957 ATP-bindi	gthLength 0 229 2-binding case (6] [LN:Al	protes sette l E001033 in] [GI	in homolog] nomology] 3] N:AF1018] gidus section 74 of
ORF Name A17503000986_35955213_c3_302 Description	NTID 1412	AAID N Len	gthLength	score	probability

ORF Name	NTID	AAID	Length Length score	probability
AT7503000986_36140963_£2_120	1413	5185		1.9e-15
Description	<u> </u>		J	
sp:[LN:TAGF_BACSU] [AC:P13485] [GN:T ACID BIOSYNTHESIS PROTEIN F] [SP:P13 [AC:S06049:G69720] [PN:probable CDP:CDP-glycerol:polyglycerol phosphate protein:teichoic-acid synthase] [GN:[EC:2.7.8.12] [DB:pir2] [MP:310 degr [OR:Bacillus subtilis] [DB:genpept-b (tag3) polypeptide (AA 1-746)] [SP:P>gp:[GI:e1184478:g2636098] [LN:BSUB0 [PN:CDP-glycerol:polyglycerol phosph [OR:Bacillus subtilis] [DB:genpept-b 19 of 21): from 3597091to 3809700.] [LE:78129] [RE:80369] [DI:complement [AC:Z99122:AL009126] [PN:CDP-glycero acid biosynthesis] [OR:Bacillus subt genome (section 19 of 21): from 3597 [SP:P13485] [LE:78129] [RE:80369] [D	485] [D glycero glycer rodC:ta ees] > ct1] [D 13485] 019] [A ate] [C] [NT:alt] > gp:[C] 1:polygilis] [C] 091to 3	B:swiss l glyce o-phosp g-3:tag gp:[GI: E:Bacil [LE:217 C:Z9912 N:tagF] E:Bacil ernate GI:e118 lycerol DB:genp 809700	sprot] >pir:[LN:S06 erophosphotransferase photransferase tagF gF] [OR:Bacillus su g40100] [LN:BSRODC] llus subtilis rodC of 78] [RE:4418] [DI:di 22:AL009126] [FN:teichoic acid llus subtilis comple gene name: rodC] [S 34478:g2636098] [LN L phosphate] [GN:tag pept] [DE:Bacillus supple	ode9] se, :rodC ubtilis]] [AC:X15200] operon.] [NT:rodC irect] biosynthesis] ete genome (section SP:P13485] :BSUB0019] gF] [FN:teichoic subtilis complete
ORF Name	NTID	AAID	NT AA LengthLength	probability
A17503000986_3906385_f3_183	1414	5186	126 41	
<u>Description</u> NO-HIT				
	and the second second section (W. was an audit of the con-		- Van
ORF Name	NTID	AAID	NT AA LengthLength	probability
AI7503000986_3957511_c2_248	1415	5187	126 41	
Description				
NO-HIT		Not consider the parties of the contract of th		
ORF Name	NTID	AAID	NT AA Length Length	probability
A17503000986_4062562_c2_269	1416	5188	183 60	
Description NO-HIT				
ORF Name	NTID	AAID	<u>NT AA</u> LengthLength	probability
A17503000986_4178140_c2_270	1417	5189	141 46	
Description No. 117				
NO-HIT	Accessed to the second		Strand Conference of the Confe	
ORF Name	NTID	AAID	NT AA LengthLength	probability
A17503000986_4297627_±2_119	1418	5190	132 43	
Description NO-HIT				
ORF Name	NTID	AAID	NT AA LengthLength	probability
A17503000986_4382062_f1_19	1419	5191	189 62	
Description				
NO-HIT				

ORF Name	NTID	AAID	NT LengthLe	AA ngth score	probability
AI7503000986_4496062_c3_287	1420	5192	345 11		2e-12
Description			J[] [
pir: [LN:E69764] [AC:E69764] [PN:hyposubtilis] [DB:pir2] >gp: [GI:e1182361 [GN:ycnI] [FN:unknown] [OR:Bacillus complete genome (section 3 of 21): fi [DI:complement] >gp: [GI:d1009660:g18 [OR:Bacillus subtilis] [SR:Bacillus [DE:Bacillus subtilis] [SR:Bacillus complete cds.] [LE:125407] [RE:12602]	:g263269 subtilis rom 4027 05463] subtilis degree 1	95] [Li 3] [DB 751 too [LN:D50 5 (stracegion	N:BSUB000 :genpept- 511850.] 0453] [AC ain:168 t containi	3] [AC:Z993 bct1] [DE:F [LE:43004] ::D50453] [C :rpC2) DNA]	106:AL009126] Bacillus subtilis [RE:43618] GN:ycnI] [DB:genpept-bct1]
ORF Name A17503000986_4687893_c1_213	NTID 1421	<u>AAID</u> 5193	NT LengthLe		probability
Description					
<pre>gp:[GI:g2822338] [LN:AF016485] [AC:A] [DB:genpept-bct2] [DE:Halobacterium sequence.] [NT:ORF H0660; similar to [DI:complement] >gp:[GI:g2822427] [L] NRC-1] [DB:genpept-bct2] [DE:Halobacsequence.] [NT:ORF H1696; similar to [DI:direct]</pre>	sp. NRC- ORF in N:AF0164 terium s	-1 plas Sulfol 185] [1 sp. NRC	smid pNRC lobus] [L AC:AF0164 C-1 plasm	100, comple E:60258] [F 85] [OR:Ha] id pNRC100,	ete plasmid RE:61397] Lobacterium sp. complete plasmid
ORF Name	NTID	AAID	<u>NT</u> LengthLe	AA ngth score	probability
AI7503000986_4689007_f1_49	1422	5194	123 40		
Description NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> LengthLe	AA ngth	probability
A17503000986_4727217_f1_25	1423	5195	402 13	3	
Description NO-HIT					
ORF Name	NTID	AAID	<u>NT</u> LengthLe	AA ngth	probability
A17503000986_4797125_f3_148	1424	5196	1647 54	8 970 1	.2e-97
Description					
sp:[LN:ARAB_BACSU] [AC:P94524] [GN:AI [DE:L-RIBULOKINASE,] [SP:P94524] [DB [PN:L-ribulokinase araB] [GN:araB] >gp:[GI:e1184128:g2635344] [LN:BSUB00 [GN:araB] [FN:L-arabinose utilization [EC:2.7.1.16] [DE:Bacillus subtilis of 3013540.] [SP:P94524] [LE:149661] [RI >gp:[GI:e1165307:g1770015] [LN:BSZ752 [OR:Bacillus subtilis] [DB:genpept-book [NT:homology to araB of Escherichia of [RE:20246] [DI:direct]	:swisspr [OR:Baci 015] [AC n] [OR:E complete E:151343 208] [AC ct1] [DE	cot] >p llus s l:Z9911 Bacillus genom [DI: l:Z7520 l:B.sub	oir:[LN:D subtilis] .8:AL0091 as subtil ne (secti- compleme .8] [PN:L otilis ge	69587] [AC: [DB:pir2] 26] [PN:L-r is] [DB:gen on 15 of 21 nt] -ribulokina nomic seque	D69587] ribulokinase] spept-bct1] ribulokinase] spept-bct1] rice [GN:araB] rice 89009bp.]

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probab	oility
A17503000986_4881588_c2_258	1425	5197	465	154	_	4.7e-09	
Description	L					•	
pir:[LN:A69783] [AC:A69783] [PN:tra [GN:ydgG]] [OR:Bacillus subtilis] [D [AC:AB001488]] [GN:ydgG]] [OR:Bacillus subtiregionbetween 35 and 47 degree.] [NT [DI:direct]] >gp:[GI:e1182530:g263286] [FN:unknown] [OR:Bacillus subtilis]] genome (section 3 of 21): from 40275 regulator (MarR family)] [LE:205898] [LN:BSUB0004] [AC:Z99107:AL009126] [DB:genpept-bct1] [DE:Bacillus subtito813890.] [NT:similar to transcript [DI:direct]	B:pir2] subtil lis gen :FUNCTI 4] [LN: [DB:gen] 1 to611: [RE:20: GN:ydgG] lis com	>gp:[is] [S ome se ON UNK BSUB00 pept-b 850.] 6356] [FN: plete	GI:d102 R:Bacil quence, NOWN.] 03] [AC ct1] [I [NT:sim [DI:dir unknown genome	20150:g llus su 148 k [LE:14 C:Z9910 DE:Baci milar t rect] > (secti	(188137 (btilis (b sequence) (2152] (6:ALOC (1lus second) (50 transpector) (50 gp:[GI (6) Bacill (6) 4 cond	(O) [LN: A) (Strain lence of [RE:1426] [G] (Subtilis lescription in the color subtilus subtilus subtilus subtilus subtilus subtilus subtilus f 21): f	AB001488] 1:168) DNA] the 510] SN:ydgG] complete onal 13:g2632877] .lis] From 600701
ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	score	probab	ility
AT7503000986_5111253_f1_6	1426	5198		48			
Description					ı		
NO-HIT							
ORF Name	NTID	AAID	NT Length	1.014	score	probab	ility
A17503000986_5128587_±1_12	1427	5199	696	231	336	1.8e-30	
Description pir:[LN:A69811] [AC:A69811] [PN:con [CL:hypothetical protein HI0278] [OR >gp:[GI:e1182755:g2633089] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-b 5 of 21): from 802821 to1011250.] [N [RE:35430] [DI:direct] >gp:[GI:d1023 [OR:Bacillus subtilis] [SR:Bacillus [DE:Bacillus subtilis 35.7 kb genomi [LE:13092] [RE:13757] [DI:complement	:Bacilli 005] [Ad ct1] [DI T:simila 167:g244 subtilis c DNA,	us sub C:Z991 E:Baci ar to 43233] s (str	tilis] 08:AL00 llus su hypothe [LN:D8 ain:AC3	[DB:pi 99126] obtilis tical 6417]	r2] [GN:yf compl protei [AC:D8 A] [DB	lK] [FN: ete geno ns] [LE: 6417] [P :genpept	unknown] me (section 34765] N:YflK] -bctl]
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probab	ility
A17503000986_5283390_c2_259	1428	5200	1416		465	3.9e-44	
Description			- L				
gp:[GI:g3676414] [LN:AF051917] [AC:A aureus] [DB:genpept-bct2] [DE:Staphy [NT:Orf423] [LE:755] [RE:2026] [DI:d	lococcus						
ORF Name	NTID	AAID	<u>NT</u> Length:	<u>AA</u> Length	score	probab	ility
A17503000986_6307_£3_182	1429	5201		57			
Description			الــــا	J			

ORF Name	MIID	AAID	Lengt	h Length	SCOLE	probabi	lity
AI7503000986_6930462_c2_260	1430	5202	1227		634	4.9e-62	
Description pir: [LN:E69783] [AC:E69783] [PN:bing) [CL:bicyclomycin resistance protections of the protection of the protect of the prote	ein] [OR:] 01488] [A(Bacill C:AB00 :genpe een 35 0] [RE C:Z991 E:Baci	us sub 1488] pt-bct and 4 :14806 07:AL0	tilis] [GN:ydo :1] [DE :7 degre :8] [DI :09126]	[DB:pgK] [O:Bacil :Bacil :ee.] [i :direc [GN:yes comp	ir2] R:Bacillus lus subti] NT:SIMILAF t] dgK] [FN:u	s subtilis] lis genome R TO unknown] me (section
ORF Name	NTID	AAID	NT Lengt	<u>AA</u> hLength	score	probabi	lity
A17503000986_7242812_f3_152	1431	5203	627	208	373	2.2e-34	
sp:[LN:3MGH_BACSU] [AC:P94378] [GN [DE:PUTATIVE 3-METHYLADENINE DNA GROWN Spir:[LN:D70082] [AC:D70082] [PN:16] [GN:yxlJ] [FN:D6] [GN:yxlJ] [FN:D6] [GN:yxlJ] [FN:D6] [GN:yxlJ] [FN:D6] [GN:yxlJ] [FN:Bacillus subtilis complete generation of the subtilis [GI:d1012408:grown subtilis] [SR:Bacillus [GR:Bacillus subtilis] [SR:Bacillus genome sequenation of the subtilis	LYCOSYLASI DNA-3-meth [DB:pir2] :unknown] ome (sect: glycosida 1783264] s subtiliance covers	E,] [Shylade >gp:[[OR:Bion 20 ase] [[LN:D8 s (string li	P:P943 nine g GI:e11 acillu of 21 SP:P94 3026] ain:BG c-cel	78] [DE 1900 1	3:swis: dase ho g26363: ilis] n 3798- LE:1640 3026:D4	sprot] omolog yxl 97] [LN:BS [DB:genper 401to 4010 671] [RE:1 45911] [GN	SUB0020] bt-bct1] 0550.] 165261] V:yxlJ] ept-bct1]
ORF Name	NTID	AAID	<u>NT</u> Lengtl	<u>AA</u> hLength	score	probabi	lity
A17503000986_803137_f1_5	1432	5204	165	54]		
Description NO-HIT							
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> nLength	score	probabi	lity
A17503000986_804837_f1_59 Description	1433	5205	963	320	342	4.3e-31	
gp:[GI:e327689:g2407930] [LN:LLPFLI protein] [OR:Lactococcus lactis] [I (strain DB1341).] [LE:464] [RE:138:	DB:genpept	t-bct1					
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> nLength	score	probabi	lity
A17503000986_814838_c1_208	1434	5206	660	219	472	7.1e-45	
Description				·			
<pre>pir:[LN:D69821] [AC:D69821] [PN:hy subtilis] [DB:pir2] >gp:[GI:e118288 [GN:yhbJ] [FN:unknown] [OR:Bacillus complete genome (section 5 of 21): [DI:direct]</pre>	39:g263322 s subtilis	23] [L s] [DB	N:BSUB :genpe	0005] [pt-bct1	AC: Z99	9108:AL009 :Bacillus	126] subtilis

ORF Name	NTID	AAID	Length Length score	probability
AI7503000986_837578_c3_326	1435	5207		7.0e-70
Description		L	J	
pir:[LN:D70179] [AC:D70179] [PN:Na+burgdorferi] [SR:, Lyme disease spir [AC:AE001165:AE000783] [PN:Na+/H+ an burgdorferi] [SR:Lyme disease spiroc (section 51 of 70) of the complete g PID:143245] [LE:6377] [RE:7726] [DI:	ochete] tiporte: hete] [] enome.]	[DB:p: r (nhac DB:genp	ir2] >gp:[GI:g268856 C-1)] [GN:BB0637] [C pept-bct2] [DE:Borre	57] [LN:AE001165] DR:Borrelia elia burgdorferi
ORF Name	NTID	AAID	NT AA LengthLength	probability
AI7503000986_8568_c3_289	1436	5208	429 142 73 0	0.017
Description				
gp:[GI:g4151243] [LN:AF063590] [AC:A [GN:mceB] [OR:Klebsiella pneumoniae] microcin E492 precursor (mceA) and m cds.] [NT:overlapping with mceA] [LE	[DB:gen	npept-k E492 in	oct2] [DE:Klebsiella nmunity protein (mce	pneumoniae
ORF Name	NTID	AAID	NT AA LengthLength	probability
AI7503000986_901515_c3_318	1437	5209	285 94 81 0	.0019
Description				
[OR:Mitochondrion Boophilus microplu [DE:Boophilus microplus cytochrome b tRNA-Leu, tRNA-Cys, and tRNA-Met gen subunit 2 (ND2) gene, partial cds,mi [LE:<1] [RE:285] [DI:direct] >gp:[GI [PN:cytochrome b apoenzyme] [GN:Cytb [SR:southern cattle tick] [DB:genpep (Cytb) gene, partialcds; tRNA-Ser, tcompletesequence; and NADH dehydrogecds,mitochondrial genes for mitochondrial genes for mitocho	apoenzyes, comptochondrigg 173123] [OR:M:t] [DE:HRNA-Leughasse subsection of the content of the	yme (Cypleteserial ge lial ge lib) [Li itochor Boophil , tRNA-	ytb) gene, partial of equence; and NADH deenes for mitochondrials [AC:AF adrion Boophilus micus microplus cytochecys, and tRNA-Met gene, partials [AC:Decenter]	ds; tRNA-Ser, chydrogenase al products.] f110612] croplus] arome b apoenzyme genes,
ORF Name	NTID	AAID	<u>NT</u> <u>AA</u> LengthLength	probability
A17503000986_968751_f2_63	1438	5210	123 40	
Description				
NO-HIT				
ORF Name	NTID	AAID	<u>NT AA</u> LengthLength	probability
AI7503000986_975261_f3_149	1439	5211	 	.022
Description		<u> </u>		
sp:[LN:YC36_GUITH] [AC:O78501] [GN:YGDE:HYPOTHETICAL 18.4 KD PROTEIN YCF. [LN:AF041468] [AC:AF041468:X14171:X6] [PN:hypothetical chloroplast RF36] [GSR:Guillardia theta] [DB:genpept-pluce:88514] [RE:88981] [DI:direct]	36] [SP: 2349:X51 GN:ycf36	078501 L511:X1 [OR:] [DB:swissprot] >g 4504:X52158:X52912: Chloroplast Guillar	p:[GI:g3603031] X56806:M7654 7] dia theta]
ORF Name	NTID	AAID	NT AA LengthLength	probability
AI7503000986_9797911_±3_139	1440	5212	126 41	
Description NO-HIT	 J		I	

ORF Name	NTID	AAID Length Length score probability
A17503000986_984628_t2_65	1441	<u>5213 [32 43 </u>
Description		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000986_990952_f3_155	1442	5214 1209 402 744 1.1e-73
Description		
[SR:Plasmid R100 (lab_host:Escherich	ia coli	:AP000342] [GN:ydhA] [OR:Plasmid R100] strain K-12) DNA] [DB:genpept] [DE:Plasmid) to 376 residues of 404 aa] [LE:37813]
ORF Name	NTID	AAID NT AA score probability
A17503000986_9944132_f1_20	1443	5215 189 62
Description NO-HIT	A	
ORF Name	NTID	AAID NT AA score probability
AI7503000987_10188258_£1_158	1444	5216 129 42
Description		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000987_10547152_£1_153	1445	5217 144 47
Description NO-HIT		
NO-NII		
ORF Name	NTID	AAID NT AA score probability LengthLength
AI7503000987_10662763_c2_764	1446	5218 1089 362 688 9.2e-68
KD PROTEIN IN MECB-GLTX INTERGENIC R [AC:S66118:D69741] [PN:conserved by hypothetical protein yacL] [OR:Bacil [LN:BAC180K] [AC:D26185] [PN:unknown (sub_species:Marburg, strain:168) DN kilobase region of replication origi >gp:[GI:e1182022:g2632356] [LN:BSUB0	EGION] pothetic lus sub [OR:Bare] [OB:Gare] [DB:Gare] [ACCO] [ACCO] [DCCO] [DCCO] [DCCO] [DCCO]	C:Z99104:AL009126] [GN:yacL] [FN:unknown] E:Bacillus subtilis complete genome (section
ORF Name	NTID	AAID NT AA score probability
AI7503000987_10667002_c1_690	1447	5219 207 68
Description NO-HIT		
NO-IIII		

ORF Name	NTID	AAID LengthLength score probability
AI7503000987_10734838_c2_819	1448	
CEREVISIAE] [SR:,BAKER'S YEAST] [EC: [SP:P32377] [DB:swissprot] >pir:[LN: decarboxylase,:protein N3427:protein cerevisiae] [EC:4.1.1.33] [DB:pir2] [AC:X97557] [PN:diphosphomevalonate cerevisiae] [SR:baker's yeast] [DB:gene.] [SP:P32377] [LE:544] [RE:1734] [LN:SCYNR043W] [AC:Z71658:Y13139] [CD: RE:RE:RE:RE:RE:RE:RE:RE:RE:RE:RE:RE:RE:R	4.1.1.3 S63374] YNR043 [MP:14R decarbo genpept- genpept- genpept- genpept- genpept- genpept- genpept- genpept- genpept-	[OR:Saccharomyces cerevisiae] [SR:baker's chromosome XIV reading frame ORF YNR043w.]
ORF Name	NTID	AAID NT AA score probability
<pre>>gp:[GI:g4155644] [LN:AE001533] [AC: [OR:Helicobacter pylori J99] [DB:gen</pre>	J99, , ; AE00153 pept-bc	5221 273 90 67 0.0041
[DB:pir2] [MP:1] >gp:[GI:g3128293]	NTID 1450 oothetica	AAID NT AA score probability 5222 1242 413 230 1.2e-16
protein] [OR:Rhodobacter capsulatus] strain SB1003, partial genome.] [LE:	[DB:gei	cal protein] [OR:Rhodobacter capsulatus] 010496] [AC:AF010496] [PN:hypothetical enpept-bct2] [DE:Rhodobacter capsulatus [RE:55613] [DI:complement]
	[DB:gei	010496] [AC:AF010496] [PN:hypothetical enpept-bct2] [DE:Rhodobacter capsulatus
ORF Name AI7503000987_110766_f1_13 Description	[DB:gen	210496] [AC:AF010496] [PN:hypothetical enpept-bct2] [DE:Rhodobacter capsulatus [RE:55613] [DI:complement] AAID NT AA LengthLength score probability

NO-HIT

ORF Name	NTID	AAID NT AA score probability				
AI7503000987_12267167_c2_786	1454	5226 135 44				
Description	L J					
NO-HIT						
ORF Name	NTID	AAID NT AA score probability				
AI7503000987_1227250_f3_539	1455	5227 996 331 562 2.1e-54				
Description	<u> </u>					
sp:[LN:YXEI_BACSU] [AC:P54948] [GN:YXEI:LP9A] [OR:BACILLUS SUBTILIS] [DE:HYPOTHETICAL 37.2 KD PROTEIN IN IDH-DEOR INTERGENIC REGION] [SP:P54948] [DB:swissprot] >pir:[LN:C70075] [AC:C70075] [PN:penicillin amidase homolog yxeI] [GN:yxeI] [CL:choloylglycine hydrolase] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1184679:g2636500] [LN:BSUB0021] [AC:Z99124:AL009126] [GN:yxeI] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 21 of 21): from 3999281to 4214814.] [NT:similar to penicillin amidase] [SP:P54948] [LE:62053] [RE:63039] [DI:complement] >gp:[GI:d1008921:g1408494] [LN:D45912] [AC:D45912] [GN:yxeI] [OR:Bacillus subtilis] [SR:Bacillus subtilis (strain:BGSC 1A1 (Marburg 168; trpC2)) DNA] [DB:genpept-bct1] [DE:Bacillus subtilis genome sequence between the iol and hut operon,partial and complete cds.] [NT:homologous to penicillin acylase] [LE:8437] [RE:9423] [DI:direct]						
ORF Name	NTID	AAID NT AA score probability				
AI7503000987 12516511 c2 813	1456	5228 129 42 164 4.7e-12				
Description						
<pre>gp:[GI:g1022726] [LN:SHU35635] [AC:U haemolyticus] [SR:Staphylococcus hae [DE:Staphylococcus haemolyticus IS12 [LE:1101] [RE:1922] [DI:complement]</pre>	molytic					
ORF Name	NTID	AAID NT AA score probability				
A17503000987_12601637_c3_898	1457	5229 144 47 76 0.0065				
<pre>disease spirochete] [DB:pir2] >gp:[G [PN:ribosomal protein S12 (rpsL)] [G disease spirochete] [DB:genpept-bct2</pre>	in S12] I:g26882 N:BB038] [DE:Bo	protein S12] [GN:BB0387:rpsL] [OR:Borrelia burgdorferi] [SR:, Lyme 295] [LN:AE001144] [AC:AE001144:AE000783] [OR:Borrelia burgdorferi] [SR:Lyme prrelia burgdorferi (section 30 of 70) of 562 percent identity: 79.03;] [LE:1809]				
ORF Name	NTID	AAID NT AA score probability				
A17503000987_12714833_c2_863	1458	5230 1218 405 95 0.021				
Description						
pir:[LN:S72278] [AC:S72278:S78483]	[PN:ATP-	-dependent Clp proteinase, homolog] [GN:clpC				

] [OR:plastid Plasmodium falciparum] [EC:3.4.21.-] [DB:pir2]

	NTID	AAID <u>NT AA</u> score probability
AI7503000987_12902217_f2_349	1459	5231 681 226 292 8.5e-26
subtilis] [DB:pir2] >gp:[GI:e1185418 [GN:yojG] [FN:unknown] [OR:Bacillus complete genome (section 11 of 21): [DI:complement] >gp:[GI:g3169323] [LifN:unknown] [OR:Bacillus subtilis] (yojA), YojB (yojB), YojC (yojC), YoYojH (yojH), YojI(yojI), YojJ (yojJ)	:g26343 subtili from 20 N:AF026 [DB:gen] jD(yojD , YojK	al protein yojG] [GN:yojG] [OR:Bacillus 39] [LN:BSUB0011] [AC:Z99114:AL009126] s] [DB:genpept-bct1] [DE:Bacillus subtilis 00171to 2207900.] [LE:120715] [RE:121143] 147] [AC:AF026147] [PN:YojG] [GN:yojG] pept-bct2] [DE:Bacillus subtilis YojA), YojE (yojE), YojF (yojF), YojG (yojG), (yojK), YojL (yojL), YojM (yojM), cds; and OdhA (odhA) gene,partial cds.]
ORF Name	NTID	AAID NT AA score probability
AI7503000987_12932802_£1_115	1460	5232 [126] [41]
Description NO-HIT		
ORF Name	NTID	AAID NT AA score probability
AI7503000987_134702_£1_113	1461	5233 834 277 673 3.6e-66
[DE:(HMP-P KINASE)] [SP:P39610] [DB: [PN:phosphomethylpyrimidine kinase to [CL:phosphomethylpyrimidine phosphate	swisspro hiD:pro	=
[DB:genpept-bct1] [DE:B.subtilis general [RE:56603] [DI:complement] >gp:[GI:e] [AC:Z99123:AL009126] [PN:phosphomether [PR:phosphomether] [OR:Bacillus subtilis]	omic req 1186301 ylpyrim: [DB:gen of 21)	N:ipa-52r] [OR:Bacillus subtilis] gion (325 to 333).] [SP:P39610] [LE:55788] :g2636337] [LN:BSUB0020] idine kinase] [GN:thiD] [FN:thiamin npept-bct1] [EC:2.7.4.7] [DE:Bacillus : from 3798401to 4010550.] [NT:alternate
[DB:genpept-bct1] [DE:B.subtilis general [RE:56603] [DI:complement] >gp:[GI:election 20] [PN:phosphometheliosynthesis] [OR:Bacillus subtilis] subtilis complete genome (section 20]	omic req 1186301 ylpyrim: [DB:gen of 21)	N:ipa-52r] [OR:Bacillus subtilis] gion (325 to 333).] [SP:P39610] [LE:55788] :g2636337] [LN:BSUB0020] idine kinase] [GN:thiD] [FN:thiamin npept-bct1] [EC:2.7.4.7] [DE:Bacillus : from 3798401to 4010550.] [NT:alternate
[DB:genpept-bct1] [DE:B.subtilis generated [RE:56603] [DI:complement] >gp:[GI:electric [AC:Z99123:AL009126] [PN:phosphomethed biosynthesis] [OR:Bacillus subtilis] subtilis complete genome (section 20 gene name: ywdB, ipa-52r] [SP:P39610] ORF Name AT7503000987_13695125_c2_771	omic req 1186301 ylpyrim: [DB:gen of 21)] [LE:10	N:ipa-52r] [OR:Bacillus subtilis] gion (325 to 333).] [SP:P39610] [LE:55788] :g2636337] [LN:BSUB0020] idine kinase] [GN:thiD] [FN:thiamin npept-bct1] [EC:2.7.4.7] [DE:Bacillus : from 3798401to 4010550.] [NT:alternate 01359] [RE:102174] [DI:direct]
[DB:genpept-bct1] [DE:B.subtilis general [RE:56603] [DI:complement] >gp:[GI:electorial [PN:phosphomether [PN:phosphometh	omic registration of 21) [LE:10] NTID [1462] S8131] E:Clostines, cor	N:ipa-52r] [OR:Bacillus subtilis] gion (325 to 333).] [SP:P39610] [LE:55788] :g2636337] [LN:BSUB0020] idine kinase] [GN:thiD] [FN:thiamin npept-bct1] [EC:2.7.4.7] [DE:Bacillus : from 3798401to 4010550.] [NT:alternate 01359] [RE:102174] [DI:direct] AAID NT AA LengthLength [S234 603 200 117 9.1e-06] [PN:SigX] [GN:sigX] [OR:Clostridium ridium acetobutylicum pho-sigX gene region, mplete cds.] [NT:proposed ECF subfamily RNA
[DB:genpept-bct1] [DE:B.subtilis general [RE:56603] [DI:complement] >gp:[GI:electorial [PN:phosphomether properties of the complete of the complete genome (section 20 gene name: ywdB, ipa-52r] [SP:P39610] ORF Name A17503000987_13695125_c2_771 Oescription gp:[GI:g3025463] [LN:CAU58131] [AC:Usacetobutylicum] [DB:genpept-bct2] [DisphoP, phoR, sigX,orf36, and orf18 general complete genome (section 20 gene name) [DB:genpept-bct2] [DisphoP, phoR, sigX,orf36, and orf18 general complete general comp	omic registration of 21) [LE:10] NTID [1462] 58131] E:Clostines, cor	N:ipa-52r] [OR:Bacillus subtilis] gion (325 to 333).] [SP:P39610] [LE:55788] :g2636337] [LN:BSUB0020] idine kinase] [GN:thiD] [FN:thiamin npept-bct1] [EC:2.7.4.7] [DE:Bacillus : from 3798401to 4010550.] [NT:alternate 01359] [RE:102174] [DI:direct] AAID NT AA LengthLength score probability [S234 603 200 117 9.1e-06 [PN:SigX] [GN:sigX] [OR:Clostridium ridium acetobutylicum pho-sigX gene region, mplete cds.] [NT:proposed ECF subfamily RNA [DI:direct]
[DB:genpept-bct1] [DE:B.subtilis generated [RE:56603] [DI:complement] >gp:[GI:electorial points of the complement] >gp:[GI:g3025463] [LN:CAU58131] [AC:Ulacetobutylicum] [DB:genpept-bct2] [Display of the complement polymerase sigmafactor] [LE:3293] [Rispandent polymerase sigmafactor] [Rispande	omic registration of 21) [LE:10] NTID [1462] 58131] E:Clostines, con E:3847]	N:ipa-52r] [OR:Bacillus subtilis] gion (325 to 333).] [SP:P39610] [LE:55788] :g2636337] [LN:BSUB0020] idine kinase] [GN:thiD] [FN:thiamin npept-bct1] [EC:2.7.4.7] [DE:Bacillus : from 3798401to 4010550.] [NT:alternate 01359] [RE:102174] [DI:direct] AAID NT AA LengthLength Score probability [PN:SigX] [GN:sigX] [OR:Clostridium ridium acetobutylicum pho-sigX gene region, mplete cds.] [NT:proposed ECF subfamily RNA [DI:direct] NT AA
[DB:genpept-bct1] [DE:B.subtilis generation [RE:56603] [DI:complement] >gp:[GI:e. [AC:Z99123:AL009126] [PN:phosphomethy biosynthesis] [OR:Bacillus subtilis] subtilis complete genome (section 20 gene name: ywdB, ipa-52r] [SP:P39610 ORF Name AT7503000987_13695125_c2_771 Description [GI:g3025463] [LN:CAU58131] [AC:Uacetobutylicum] [DB:genpept-bct2] [DisphoP, phoR, sigX,orf36, and orf18 generation [LE:3293] [RIDEF Name	omic registration of 21) [LE:10] NTID [1462] 58131] E:Clostines, con E:3847]	N:ipa-52r] [OR:Bacillus subtilis] gion (325 to 333).] [SP:P39610] [LE:55788] :g2636337] [LN:BSUB0020] idine kinase] [GN:thiD] [FN:thiamin npept-bct1] [EC:2.7.4.7] [DE:Bacillus : from 3798401to 4010550.] [NT:alternate 01359] [RE:102174] [DI:direct] AAID NT AA Score probability [EX:334 603 200 117 9.1e-06 [PN:SigX] [GN:sigX] [OR:Clostridium ridium acetobutylicum pho-sigX gene region, mplete cds.] [NT:proposed ECF subfamily RNA [DI:direct] AAID NT AA LengthLength Score probability [AAID NT AA LengthLength Score probability
[DB:genpept-bct1] [DE:B.subtilis generation [RE:56603] [DI:complement] >gp:[GI:e. [AC:Z99123:AL009126] [PN:phosphomethy biosynthesis] [OR:Bacillus subtilis] subtilis complete genome (section 20 gene name: ywdB, ipa-52r] [SP:P39610 ORF Name AT7503000987_13695125_c2_771 Description gp:[GI:g3025463] [LN:CAU58131] [AC:U.acetobutylicum] [DB:genpept-bct2] [DisphoP, phoR, sigX,orf36, and orf18 generation polymerase sigmafactor] [LE:3293] [RIDORF Name AT7503000987_1379061_c3_991 Description	omic registration of 21) [LE:10] NTID [1462] 58131] E:Clostines, con E:3847]	N:ipa-52r] [OR:Bacillus subtilis] gion (325 to 333).] [SP:P39610] [LE:55788] :g2636337] [LN:BSUB0020] idine kinase] [GN:thiD] [FN:thiamin npept-bct1] [EC:2.7.4.7] [DE:Bacillus : from 3798401to 4010550.] [NT:alternate 01359] [RE:102174] [DI:direct] AAID NT AA Score probability [EX:334 603 200 117 9.1e-06 [PN:SigX] [GN:sigX] [OR:Clostridium ridium acetobutylicum pho-sigX gene region, mplete cds.] [NT:proposed ECF subfamily RNA [DI:direct] AAID NT AA LengthLength Score probability [AAID NT AA LengthLength Score probability
[DB:genpept-bct1] [DE:B.subtilis generation [RE:56603] [DI:complement] >gp:[GI:electorial subtilis] [AC:Z99123:AL009126] [PN:phosphomether biosynthesis] [OR:Bacillus subtilis] subtilis complete genome (section 20 gene name: ywdB, ipa-52r] [SP:P39610] ORF Name A17503000987_13695125_c2_771 Description gp:[GI:g3025463] [LN:CAU58131] [AC:Usacetobutylicum] [DB:genpept-bct2] [DisphoP, phoR, sigX,orf36, and orf18 genpept-book polymerase sigmafactor] [LE:3293] [RIDDER Name A17503000987_1379061_c3_991 Description NO-HIT	omic registration of 21) [DB:gen of 21) [LE:10] NTID [1462] 58131] E:Clostman context of 21 NTID [1463] NTID	N:ipa-52r] [OR:Bacillus subtilis] gion (325 to 333).] [SP:P39610] [LE:55788] :g2636337] [LN:BSUB0020] idine kinase] [GN:thiD] [FN:thiamin npept-bct1] [EC:2.7.4.7] [DE:Bacillus : from 3798401to 4010550.] [NT:alternate 01359] [RE:102174] [DI:direct] AAID NT AA Score probability [PN:SigX] [GN:sigX] [OR:Clostridium ridium acetobutylicum pho-sigX gene region, mplete cds.] [NT:proposed ECF subfamily RNA [DI:direct] AAID NT AA Score probability 5235 153 50

ORF Name

ORF Name	NTID	AAID LengthLength score probability
AI7503000987_13876005_c1_677	1465	
Description		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
		Length Length
A17503000987_14254437_c3_903	1466	5238 123 40
Description NO-HIT		
ORF Name	NTID	AAID LengthLength score probability
A17503000987_14460882_f1_107	1467	5239 177 58 236 7.3e-20
Description		
gp:[GI:g1022726] [LN:SHU35635] [AC:U		
haemolyticus] [SR:Staphylococcus hae		us strain=Y176] [DB:genpept-bct1] and ORF2 genes, completecds.] [NT:ORF1]
[LE:1101] [RE:1922] [DI:complement]	1/2 ORF1	and ORF2 genes, completecds.] [NT:ORF1]
		NT AA
ORF Name	NTID	AAID LengthLength score probability
A17503000987_14501556_f2_325	1468	5240 [171] [56
Description		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000987 14537578 c2 763	1469	Length Length Score Plobability [5241 [147 [48]
Description	1405	3241 147 40
NO-HIT		
		NITT 7.7
ORF Name	NTID	AAID LengthLength score probability
A17503000987_14547143_c3_912	1470	5242 522 173 501 6.0e-48
Description		
		R:BACILLUS SUBTILIS] [DE:HYPOTHETICAL 17.8
[AC:S11690:S66048:B69737] [PN:conse		[SP:P21335] [DB:swissprot] >pir:[LN:S11690]
[CL:hypothetical protein Yaaj] [OR:B	acillus	subtilis] [DB:pir2]
		D26185] [PN:unknown] [OR:Bacillus subtilis]
		strain:168) DNA] [DB:genpept-bct1] [DE:B. ation origin.] [LE:89452] [RE:89937]
[DI:direct] >gp:[GI:g40011] [LN:BSOR		
		for ORF17, small cytoplasmic RNA and
		P:P21335] [LE:129] [RE:614] [DI:direct] C:Z99104:AL009126] [GN:yaaJ] [FN:unknown]
		E:Bacillus subtilis complete genome (section
1 of 21): from 1 to213080.] [NT:simi		
[LE:25850] [RE:26335] [DI:direct]		
ORF Name	NTID	AAID NT AA score probability
		AAID LengthLength score probability
A17503000987_14587817_f3_520	1471	5243 528 175
Description NO-HIT		
MO - III I		

ORF Name	NTID	AAID	NT AA score probability
A17503000987_14647510_f1_27	1472	5244	255 84
Description NO-HIT			
NO-HII			
ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000987_14879688_c1_704	1473	5245	927 308 234 5.1e-26
Description			
pir:[LN:A71042] [AC:A71042] [PN:pro [OR:Pyrococcus horikoshii] [DB:pir2] [AC:AP000006:AB005215:AB009510:AB009 hypothetical mevalonate kinase] [GN:horikoshii (strain:OT3) DNA, clone:P [DE:Pyrococcus horikoshii OT3 genomi [NT:similar to owl:MTU47134 percent [DI:complement]	>gp:[G] 511:AB00 PH1625] yrococcu c DNA, 1	:d103 9512: [OR:P is hor	1680:g3258054] [LN:AP000006] AB009513:AB009514] [PN:335aa long yrococcus horikoshii] [SR:Pyrococcus ikoshi] [DB:genpept-bct1] 1-1485000 nt. position(6/7).]
ORF Name	NTID	AAID	NT AA ggore probability
			Length Length probability
A17503000987_14882681_c1_717	1474	5246	210 69 99 2.4e-05
Description gp:[GI:g2897106] [LN:AF020798] [AC:A			
thermophilus bacteriophage TP-J34] [bacteriophage lysogeny module, integ surface-exposed lipoprotein, putative regulatoryprotein, and P1-antirepres regulatory protein; orf121] [LE:4825	rasehomo metallo sor homo	olog (o-protolog g	int), putative host cell einase, repressor, Cro-like enes, complete cds.] [NT:CI-like
ORF Name	NTID	AAID	NT AA score probability
A17503000987_14885260_c2_839	1475	5247	441 146 209 5.3e-17
Description gp:[GI:g2444132] [LN:U88974] [AC:U88 temperate bacteriophage O1205] [DB:g temperate bacteriophage O1205, compl terminase] [LE:13835] [RE:14329] [DI	enpept-p etegenom	hg] [1 1e.] [1	DE:Streptococcus thermophilus
ORF Name	NTID	AAID	NT AA score probability
AI7503000987_15735181_f3_451	1476	5248	204 67
Description		L	
NO-HIT			
ODE Name		3375	NT AA
ORF Name	NTID	AAID	LengthLength score probability
AI7503000987_15782160_c3_905	1477	5249	747 248 469 1.5e-44
Description			_
<pre>gp:[GI:g1458327] [LN:CELF08F3] [AC:U6 [SR:Caenorhabditis elegans strain=Br2 elegans cosmid F08F3.] [LE:1515:1817 [DI:directJoin]</pre>	istol N2] [DB	genpept-inv1] [DE:Caenorhabditis:

ORF Name A17503000987_15892932_c3_902 Description NO-HIT	NTID 1478	 	NT AA score LengthLength 129 42	probability
ORF Name AI7503000987_16054827_c3_959 Description pir:[LN:T00183] [AC:T00183] [PN:hyp phage phi PVL] [DB:pir3] >gp:[GI:d10 [OR:bacteriophage phi PVL] [SR:bacte aureus ATC] [DB:genpept-phg] [DE:Bac sequence.] [NT:orf 50] [LE:36701] [R	32884:g: riophage teriopha	5251 al prote 3341957; e phi Porte age phi	ein 50] [OR:Staphyl] [LN:AB009866] [AC VL (specific_host:S PVL proviral DNA,	:AB009866] taphylococcus
ORF Name A17503000987_16212803_c2_844 Description pir: [LN:S58144] [AC:S58144] [PN:gen >gp: [GI:e244843:g2764862] [LN:BSPP1G [DB:genpept-phg] [DE:Bacteriophage S [LE:9012] [RE:9320] [DI:direct] >gp: [OR:Bacteriophage SPP1] [DB:genpept-genes 7 to 15.] [NT:product required [DI:direct]	ENM] [AG PP1 comp [GI:g10! phg] [DI	5252 otein] C:X97918 olete no 52813] E:Bacter	[OR:phage SPP1] [DB B] [OR:Bacteriophage ucleotide sequence. [LN:SPP1HEADG] [AC: riophage SPP1 head	e SPP1]] [NT:gene 15] X89721] [GN:15] morphogenesis
ORF Name A17503000987_16413130_f1_128 Description gp:[GI:g3929312] [LN:AF100426] [AC:A [GN:fap1] [OR:Streptococcus parasang parasanguis fimbriae-associated prot fimbriae assembly and fimbriae-media	uinis] ein Fap1	5253 [PN:fi] [DB:genr	imbriae-associated poept-bct2] [DE:Strept	ptococcus .] [NT:invovled in
ORF Name A17503000987_16603427_c2_861 Description NO-HIT	NTID 1482		NT AA engthLength 531 176	probability
ORF Name A17503000987_16681687_c1_635 Description sp:[LN:YACI_BACSU] [AC:P37570] [GN:Y. KD PROTEIN IN LYSS-MECB INTERGENIC R. >pir:[LN:S66114] [AC:S66114:I40507:B] [OR:Bacillus subtilis] [DB:pir2] > [PN:unknown] [OR:Bacillus subtilis] strain:168) DNA] [DB:genpept-bct1] [Preplication origin.] [LE:166083] [RE [LN:BSUB0001] [AC:Z99104:AL009126] [OB:genpept-bct1] [DE:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis] [DI:direct]	EGION (0 69741] gp:[GI:0 [SR:Baci DE:B. su :167174] GN:yacI] lis comp	E:BACILI (PN:credinos861) (Incredinos861) (Incredinos861) (Incredinos961) (Incredinos961) (Incredinos961)	LUS SUBTILIS] [DE:HI [SP:P37570] [DB:swisted in the complete of the complete	ssprot] og yacI] [GN:yacI BOK] [AC:D26185] s:Marburg, region of 2018:g2632352] s subtilis] 21): from 1

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000987_16798777_c1_646	1484	5256	3558	1185	5906	0.0
Description sp:[LN:RPOB_STAAU] [AC:P47768] [GN:R] [DE:BETA CHAIN) (RNA POLYMERASE BETA >pir:[LN:S59951] [AC:S59951] [PN:DN] [CL:DNA-directed RNA polymerase beta [DB:pir2] >gp:[GI:g677851] [LN:SARPL] beta chain] [GN:rpoB] [OR:Staphylocod [DE:S.aureus rplL, orf202, rpoB(rif) hypothetical protein ORF202, DNA-dire [SP:P47768] [LE:1222] [RE:4770] [DI:comparison of the comparison of the compa	SUBUNITA-direct chain] RPO] [ACccus aur and rpo	c)] [SI ced RNA [OR:St C:X641] ceus] oC gene	P:P4776 A polym caphylo 72] [PN [DB:gen es for	[DB] nerase, neoccus l:DNA-d npept-b riboso	beta aureu irecte ctl]	sprot] chain] [GN:rpoB] us] [EC:2.7.7.6] ed RNA polymerase [EC:2.7.7.6] btein L7/L12,
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000987_16972575_f1_196	1485	5257	186	61	76	0.036
Description gp:[GI:g4580755] [LN:AF061085] [AC:Al herbaceum] [DB:genpept-pln2] [DE:Goss [NT:similar to P-glycoprotein in Horo	sypium h	erbace	eum P-g	lycopr	otein	gene, partial cds.]
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000987_1702_£2_209	1486	5258	522	173	88	0.0089
pir:[LN:T00168] [AC:T00168] [PN:hypophage phi PVL] [DB:pir3] >gp:[GI:d103] [OR:bacteriophage phi PVL] [SR:bacteriophage phi PVL] [DB:genpept-phg] [DE:Bacteriophage] [DE:Bacteriophage] [NT:orf 33] [LE:28172] [RI	32869:g3 riophage ceriopha	341942 phi I ge phi	?) [LN: PVL (sp L PVL p	AB0098 ecific rovira	66] [<i>A</i> _host:	AC:AB009866] Staphylococcus
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000987_19690876_c3_935	1487	5259	876	291	662	5.3e-65
Description sp:[LN:YWFL_BACSU] [AC:P39648] [GN:YW [DE:HYPOTHETICAL 31.4 KD PROTEIN IN IMPORTED IN IMPORT IN IMP	PTA 3'RE [PN:ywf 3:pir2] [DB:genp [RE:94 GN:ywfL] Lis comp	GION] L prot >gp:[G ept-bc 145] [FN:u	[SP:P3 cein:hy SI:g414 ctl] [D [DI:dir unknown genome	9648] pothet 014] [E:B.su ect] > [OR: (section	[DB:swical p LN:BSG btilis gp:[GI Bacill on 20	rissprot] protein ipa-90d] ENR] [AC:X73124] genomic region cel186264:g2636300] us subtilis] of 21): from
ORF Name	NTID	AAID	<u>NT</u> Length:	<u>AA</u> Length	score	probability
A17503000987_19695386_c3_953	1488	5260	771	256	111	0.00059
Description pir:[LN:T00180] [AC:T00180] [PN:hypo phage phi PVL] [DB:pir3] >gp:[GI:d103 [OR:bacteriophage phi PVL] [SR:bacter aureus ATC] [DB:genpept-phg] [DE:Bact sequence.] [NT:orf 46] [LE:35157] [RE	32881:g3 riophage eriopha	341954 phi P ge phi	LN: PVL (sp PVL p	AB00980 ecific	66] [Ā _host:	C:AB009866] Staphylococcus

ORF Name AAID NTID score probability LengthLength A17503000987_20078287_£3_533 1489 5261 825 2.5e-07 Description pir:[LN:B70798] [AC:B70798] [PN:probable membrane protein] [GN:Rv3737] [OR:Mycobacterium tuberculosis] [DB:pir2] >qp:[GI:e1264597:q2960161] [LN:MTV025] [AC:AL022121:AL123456] [PN:hypothetical protein Rv3737] [GN:Rv3737] [OR:Mycobacterium tuberculosis [DB:genpept-bct1] [DE:Mycobacterium tuberculosis H37Rv complete genome; segment 155/162.] [NT:Rv3737, (MTV025.085), len: 529. Probable membrane] [LE:94796] [RE:96385] [DI:direct] >gp:[GI:e1264597:g2960161] [LN:MTV025] [AC:AL022121:AL123456] [PN:hypothetical protein Rv3737] [GN:Rv3737] [OR:Mycobacterium tuberculosis] [DB:genpept] [DE:Mycobacterium tuberculosis H37Rv complete genome; segment 155/162.] [NT:Rv3737, (MTV025.085), len: 529. Probable membrane] [LE:94796] [RE:96385] [DI:direct] NT AΑ ORF Name AAID NTID score probability LengthLength AI7503000987 20087752 f2 352 1490 5262 1506 501 Description sp:[LN:TAGE BACSU] [AC:P13484] [GN:TAGE:RODD:GTAA] [OR:BACILLUS SUBTILIS] [EC:2.4.1.52] [DE: (EC 2.4.1.52) (TEICHOIC ACID BIOSYNTHESIS PROTEIN E)] [SP:P13484] [DB:swissprot] >pir:[LN:S06048] [AC:S06048:F69720] [PN:poly(glycerol-phosphate) alpha-glucosyltransferase, tagE:probable rodD protein:UDP-glucose--polyglycerol phosphate glucosyltransferase tagE] [GN:tagE:rodD] [OR:Bacillus subtilis] [EC: 2.4.1.52] [DB:pir2] [MP:310 degrees] >gp:[GI:g580920] [LN:BSRODC] [AC:X15200] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis rodC operon.] [NT:rodD (gtaA) polypeptide (AA 1-673)] [SP:P13484] [LE:157] [RE:2178] [DI:direct] >gp:[GI:e1184479:g2636099] [LN:BSUB0019] [AC:Z99122:AL009126] [PN:UDP-glucose:polyglycerol phosphate] [GN:tagE] [FN:teichoic acid biosynthesis] [OR:Bacillus subtilis] [DB:genpept-bct1] [EC:2.4.1.52] [DE:Bacillus subtilis complete genome (section 19 of 21): from 3597091to 3809700.] [NT:alternate gene name: rodD, gtaA, gtaD] [SP:P13484] [LE:80369] [RE:82390] [DI:complement] >qp: [GI:e1184479:q2636099] [LN:BSUB0019] [AC:Z99122:AL009126] [PN:UDP-glucose:polyglycerol phosphate] [GN:tagE] [FN:teichoic acid biosynthesis] [OR:Bacillus subtilis] [DB:genpept] [EC:2.4.1.52] [DE:Bacillus subtilis complete genome (section 19 of 21): from 3597091to 3809700.] [NT:alternate gene name: rodD, gtaA, gtaD] [SP:P13484] [LE:80369] [RE:82390] [DI:complement] NT AAORF Name NTID AAID score probability LengthLength AI7503000987_20322153_c2_818 1491 5263 219 72 Description NO-HIT NT AΑ ORF Name NTID AAID score probability LengthLength AI7503000987_20331552_c1_659 1492 996 5264 331 3.2e-42 Description pir:[LN:C69066] [AC:C69066] [PN:ornithine cyclodeaminase] [GN:MTH1495] [OR:Methanobacterium thermoautotrophicum] [DB:pir2] >gp:[GI:g2622612] [LN:AE000910] [AC:AE000910:AE000666] [PN:ornithine cyclodeaminase] [GN:MTH1495] [OR:Methanobacterium

thermoautotrophicum] [DB:genpept-bct2] [DE:Methanobacterium thermoautotrophicum from bases 1349621 to 1362200(section 116 of 148) of the complete genome.] [NT:Function

Code:5.09 - L-Amino Acid Metabolism,] [LE:963] [RE:1982] [DI:complement]

ORF Name	NTID	AAID	NT LengthL	AA ength S	core	probabi.	lity
A17503000987_20511590_c3_893	1493	5265			114	1.0e-38	
Description			JL L				
sp:[LN:RL7_MICLU] [AC:P02395] [GN:RPI LYSODEIKTICUS] [DE:50S RIBOSOMAL PROT >pir:[LN:R7MCML] [AC:A02771] [PN:rik [CL:Escherichia coli ribosomal protes lysodeikticus] [DB:pir1]	TEIN L7, posomal	/L12 (i prote:	MA1/MA2) in L7/L1] [SP::	P0239 somal	[DB:sw. protein l	issprot] MA]
ORF Name	NTID	AAID	<u>NT</u> LengthL	AA ength	core	probabil	lity
AI7503000987_20756260_c1_729	1494	5266	408	135	84	2.4e-14	
<u>Description</u> pir:[LN:T00194] [AC:T00194] [PN:hypo phage phi PVL] [DB:pir3] >gp:[GI:d103 [OR:bacteriophage phi PVL] [SR:bacter	32895:g3	3341968	3] [LN:A	B00986	6] [A	C:AB00986	6]
aureus ATC] [DB:genpept-phg] [DE:Bact sequence.] [NT:orf 61] [LE:39932] [RE	eriopha	age phi	i PVL pr				Jeeus
ORF Name	NTID	AAID	<u>NT</u> LengthL	<u>AA</u> ength	core	probabil	lity
AI7503000987_20980262_c2_762	1495	5267	2472	323	959	0.0	
Description sp:[LN:MECB_BACSU] [AC:P37571] [GN:MEREGULATOR OF GENETIC COMPETENCE MECB] [AC:I40508:S66115:I40385:H69600] [PN clpC:adenosine triphosphatase clpC:cl [CL:ATP-dependent Clp proteinase chai >gp:[GI:d1005862:g467474] [LN:BAC180K subtilis] [SR:Bacillus subtilis (sub [DB:genpept-bct1] [DE:B. subtilis DNF [LE:167171] [RE:169603] [DI:direct] > adenosine triphosphatase] [GN:mecB] [OR:Bacillus subtilis] [DB:genpept-bc adenosine triphosphatase (mecB)gene, [LE:335] [RE:2767] [DI:direct] >gp:[GAC:Z99104:AL009126] [PN:class III st [FN:negative regulator of late compet [DB:genpept-bct1] [DE:Bacillus subtil to213080.] [NT:alternate gene name: m [DI:direct]	[SP:P3 J:class pA/clpF n A] [G] [AC:I species A, 180 } gp:[GI: FN:comp ct1] [DF complet GI:e1182 ress re ence ge is comp	III standard	(DB:swi cress re cin homo illus su [PN:cl urg, str se regio [CO] [LN: e gene r llus sub orfX a 2632353] e-relate [OR:Bac genome (esponse- esponse- esponse- esponse- esponse- esponse- esponse- en of re BSU0260 espresse etilis N and orf: [LN:BS ed ATPas eillus s section	>pi -rela GN:cl [DB B fam B) DN eplic O4] [Or; r Marbu Y, pa SUB00 Se] [Subti	r:[LN:I40! ted ATPase pC:mecB] :pir2] ily] [OR:EA] ation orig AC:U02604] equired for rg 168 Cly rtial cds: 01] GN:clpC] lis] f 21): fro	Sacillus Gin.] [PN:ClpC or cell] oC
ORF Name A17503000987_2117202_c2_831 Description NO-HIT	NTID 1496	<u>AAID</u> 5268	NT LengthL	AA ength 8	core	probabil	ity
ORF Name AI7503000987_21484465_c2_848 Description gp:[GI:g3702331] [LN:ATAC005397] [AC:[SR:thale cress] [DB:genpept-pln2] [Dgenomic sequence.]	E:Arabi [NT:hyp	5269 97] [GN dopsis	J:T3F17. thalia cal pro	71 9 17] [OF na chro tein]	IL R:Aral	me II BAC	chaliana)

ORF Name	NTID	AAID NT AA score probability
AI7503000987_21598838_t3_534	1498	5270 465 154 155 7.2e-11
Description	·	
PROTEIN HI0108] [SP:P44520] [DB:swis [PN:hypothetical protein HI0108] [OF >gp:[GI:g1573061] [LN:U32696] [AC:U3 [GN:HI0108] [OR:Haemophilus influenz	ssprot] R:Haemop B2696:L4 Zae Rd] e genome	hilus influenzae] [DB:pir2] 2023] [PN:conserved hypothetical protein] [DB:genpept-bct2] [DE:Haemophilus influenzae .] [NT:similar to GB:U14003 SP:P39402
ORF Name	NTID	AAID NT AA score probability
AI7503000987_21603777_f1_144	1499	5271 126 41
Description NO-HIT		
ORF Name	NTID	AAID <u>NT AA</u> score probability
A17503000987_21758468_c3_992	1500	5272 489 162 123 6.9e-08
] [CL:Escherichia coli peptide N-ace [DB:pir2] >gp:[GI:g2983204] [LN:AE00 [PN:ribosomal-protein-alanine acetyl	etyltran 00696] [.transfe	<u> </u>
ORF Name	NTID	AAID NT AA score probability
AI7503000987_22042128_f3_519	1501	5273 150 49
Description NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000987_22069160_c1_652	1502	5274 393 130 623 7.1e-61
PROTEIN S12] [SP:P48942] [DB:swisspr [PN:ribosomal protein S12] [GN:rpsL]	ot] >gp [OR:Staprotein	R:STAPHYLOCOCCUS AUREUS] [DE:30S RIBOSOMAL :[GI:g706921] [LN:SAU20869] [AC:U20869] aphylococcus aureus] [DB:genpept-bct2] S12 (rpsL) gene, completecds, ribosomal cds.] [LE:418] [RE:831] [DI:direct]
ORF Name	NTID	AAID NT AA score probability
AI7503000987_2211036_f2_414	1503	5275 174 57
Description NO-HIT		
ORF Name	NTID	AAID NT AA score probability
AI7503000987_22272200_c1_735	1504	[5276 [939]312
Description NO-HIT		

ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	score	probability
AI7503000987_22400261_c2_843	1505	5277	834	277	634	4.9e-62
Description gp:[GI:e139438:g1369939] [LN:BTP9011 [OR:Bacteriophage B1] [DB:genpept-ph [LE:360] [RE:1181] [DI:direct]						
ORF Name	NTID	AAID	<u>NT</u> Lengthl	<u>AA</u> Length	score	probability
Description NO-HIT	1506	5278		42		
ORF Name	NTID	AAID	NT	<u>AA</u>	score	probability
AI7503000987_22539186_f1_67	1507	5279	Length I	45		
Description NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> LengthI	<u>AA</u> Length	score	probability
A17503000987_22689067_f1_205	1508	5280	1221	406	1218	5.3e-124
sp:[LN:NUPC_BACSU] [AC:P39141] [GN:N NUCLEOSIDE TRANSPORT PROTEIN] [SP:P3 [LN:D45912] [AC:D45912] [PN:pyrimidi [OR:Bacillus subtilis] [SR:Bacillus DNA] [DB:genpept-bct1] [DE:Bacillus operon,partial and complete cds.] [L	9141] [1 ne nucle subtilia subtilia	DB:swi eoside s (str s geno	ssprot] transpo ain:BGSO me sequo	>gp:[0 ort pro C 1A1 ence be	GI:d100 otein] (Marbu: etween	08934:g1408507] [GN:nupC] rg 168; trpC2))
ORF Name	NTID	AAID	<u>NT</u> LengthI	<u>AA</u> Length	score	probability
A17503000987_22694377_c2_776	1509	5281	741	246	762]	l.3e-75
Description pir: [LN:S59955] [AC:S59955] [PN:hyp MJ0882] [OR:Staphylococcus aureus] [[PN:hypothetical protein] [GN:ORF202 [DE:S.aureus rplL, orf202, rpoB(rif) hypothetical protein ORF202, DNA-dir [RE:1007] [DI:direct]	DB:pir2]] [OR:St and rpc] >gp: taphyl oC gen	[GI:g67 ococcus es for	7850] aureus ribosom	[LN:SAI s] [DB malprot	RPLRPO] [AC:X64172] :genpept-bct1] tein L7/L12,
ORF Name	NTID	AAID	NT LengthI	<u>AA</u> Length	score	probability
A17503000987_22773302_c1_741	1510	5282	1803	600	1371 3	3.9e-140
Description gp:[GI:e286568:g2764983] [LN:BP187PL [GN:ply187] [OR:Staphylococcus phage ply187 and hol187 genes.] [LE:222] [187] [I	DB:gen	pept-pho		_	-
ORF Name	NTID	AAID	NT LengthI	AA length	score	probability
A17503000987_22790941_c2_847	1511	5283			143 5	0.2e-10
Description gp:[GI:e244714:g2764866] [LN:BSPP1GE [DB:genpept-phg] [DE:Bacteriophage S [LE:10481] [RE:11014] [DI:direct]						

ORF Name	NTID	AAID	NT AA score	probability
AI7503000987_23442135_c1_685	1512	5284	675 224 156	7.9e-11
Description		L	ـــــا لـــــا	
pir:[LN:F71309] [AC:F71309] [PN:pro] [GN:TP0554] [OR:Treponema pallidum : [DB:pir2] >gp:[GI:g3322848] [LN:AE00] phosphatase (gph-2)] [GN:TP0554] [OR [DE:Treponema pallidum section 47 of GB:L42023 SP:P44755 PID:1004013] [LE	subsp. p 1231] [<i>I</i> :Trepone 87 of t	pallid AC:AE0 ema pai the con	um] [SR:, syphilis 01231:AE000520] [PN llidum] [DB:genpept mplete genome.] [N]	spirochete] J:phosphoglycolate -bct2]
ORF Name	NTID	AAID	NT AA LengthLength score	probability
AI7503000987_23469213_c2_838	1513	5285	411 136	
Description		——	JI	
NO-HIT				
ORF Name	NTID	AAID	NT AA LengthLength	probability
AI7503000987_23477213_c1_701	1514	5286	1011 336 1085	7.9e-110
<u>Description</u> pir:[LN:S39743] [AC:S39743:D69683]				
[CL:phosphate acetyltransferase] [OR [LN:BSGENR] [AC:X73124] [GN:ipa-88d] [DE:B.subtilis genomic region (325 to [DI:direct] >gp:[GI:e1186266:g2636302 [PN:phosphotransacetylase] [GN:pta] [EC:2.3.1.8] [DE:Bacillus subtilis co 4010550.] [NT:alternate gene name: ip [DI:complement]	[OR:Bac o 333).] 2] [LN:E [OR:Baci omplete	:illus [SP:I SSUB002 :llus s genome	subtilis] [DB:genp P39646] [LE:91234] 20] [AC:Z99123:AL00 subtilis] [DB:genpe e (section 20 of 21	ept-bct1] [RE:92205] 9126] pt-bct1]): from 3798401to
ORF Name	NTID	AAID	NT AA Length Length score	probability
A17503000987_23554700_c3_1002	1515	5287	162 53	
Description				
NO-HIT				9.00
ORF Name	NTID	AAID	NT <u>AA</u> LengthLength	probability
A17503000987_23572178_c3_958	1516	5288	195 64	
Description				
NO-HIT	NA WARRANCE TO STORY A STATE OF THE PERSON OF	AND THE RESIDENCE OF THE PARTY		
ORF Name	NTID	AAID	NT AA LengthLength	probability
A17503000987_23601637_c2_794	1517	5289	816 271 374	1.7e-34
Description				
pir:[LN:S60902] [AC:S60902:S49238:S44 [OR:Haemophilus influenzae] [DB:pir2] [OR:Haemophilus influenzae] [DB:genpe capsulation locus.] [NT:orf1] [LE:434	>gp:[G ept-bct1	I:g471] [DE:	.234] [LN:HISBCAL] H.influenzae DNA f	[AC:X78559]

ORF Name	NTID AAID LengthLength score probability
AI7503000987_23603391_c3_969	1518
[GN:F28A21.150] [OR:Arabidopsis thal	[AC:AL035526] [PN:hypothetical protein] [iana] [SR:thale cress] [DB:genpept-pln1] [some 4, BAC clone F28A21 (ESSAproject).] [26:63337] [DI:complementJoin]
ORF Name	NTID AAID NT AA score probability
A17503000987_23617140_c1_686	1519 5291 1404 467 441 6.0e-44
[GN:yhfT] [CL:acetateCoA ligase h >gp:[GI:e1183038:g2633372] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-b 6 of 21): from 999501 to1209940.] [N [LE:110451] [RE:111890] [DI:compleme [AC:Y14084] [PN:hypothetical protein [DB:genpept-bct1] [DE:B.subtilis chr	g-chain fatty-acid-CoA ligase homolog yhfT] lomology] [OR:Bacillus subtilis] [DB:pir2] 1006] [AC:Z99109:AL009126] [GN:yhfT] [FN:unknown] 10ctl] [DE:Bacillus subtilis complete genome (section of the subtilis complete genome (section of the subtiliar to long-chain fatty-acid-CoA ligase] 11. [SI:e324999:g2226256] [LN:BSY14084] 12. [GN:yhfT] [OR:Bacillus subtilis] 13. [GN:yhfT] [OR:Bacillus subtilis] 14. [GN:yhfT] [OR:Bacillus subtilis] 15. [GN:yhfT] [OR:Bacillus subtilis] 16. [GN:yhfT] [OR:Bacillus subtilis] 17. [GN:yhfT] [OR:Bacillus subtilis] 18. [GN:yhfT] [OR:Bacillus subtilis] 19. [GN:yhfT] [OR:Bacillus subtilis] 19. [GN:yhfT] [OR:Bacillus subtilis]
	NTID AAID LengthLength score probability 1520 5292 198 65 64 0.0061 F078787] [GN:T17A3.9] [OR:Caenorhabditis elegans] elegans cosmid T17A3.] [LE:15875:16120:16852] atJoin]
ORF Name	NTID AAID NT AA score probability
A17503000987_23709631_c1_676 Description NO-HIT	1521 5293 123 40
ORF Name	NTID AAID NT AA score probability
AI7503000987_23712830_c1_713	1522 5294 489 162 227 6.6e-19
bacteriophage Sfi19] [DB:genpept-phg	NT AA
ORF Name	NTID AAID LengthLength score probability
A17503000987_23867125_f2_333	1523 5295 204 67 57 0.018
Description pir:[LN:S21443] [AC:S21443] [PN:hypo [DB:pir2]	othetical protein] [OR:Dictyostelium discoideum]

ORF Name	NTID	AAID LengthLength score probability
A17503000987_23910052_f3_585	1524	5296 147 48
Description		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
AI7503000987_2392837_c1_732	1525	5297 426 141 226 8.4e-19
Description		
<pre>gp:[GI:e244844:g2764864] [LN:BSPP1GE [DB:genpept-phg] [DE:Bacteriophage S [LE:9632] [RE:10066] [DI:direct]</pre>		:X97918] [OR:Bacteriophage SPP1] plete nucleotide sequence.] [NT:gene 16.1]
ORF Name	NTID	AAID NT AA score probability
AI7503000987_24026077_c3_966	1526	5298 954 317 560 3.4e-54
<pre>>gp:[GI:e244838:g2764848] [LN:BSPP1G [DB:genpept-phg] [DE:Bacteriophage S [LE:3802] [RE:4728] [DI:direct] >gp:</pre>	ENM] [AGPP1 comp [GI:g10! phg] [DI	plete nucleotide sequence.] [NT:gene 7] 52806] [LN:SPP1HEADG] [AC:X89721] [GN:7] E:Bacteriophage SPP1 head morphogenesis
ORF Name A17503000987_24026576_c2_854 Description	NTID 1527	AAID NT AA score probability 5299 180 59
NO-HIT		
ORF Name	NTID	AAID <u>NT AA</u> score probability
AI7503000987_24229837_c1_716	1528	5300 243 80 97 3.9e-05
[OR:Lactobacillus sake] [DB:pir2] >g	p:[GI:g	al protein (bacteriocin saiA 3'-region)] 599850] [LN:LSSAKACLU] [AC:Z46867] [GN:orf1] [DE:L.sake sakacin A gene cluster.] [LE:5]
ORF Name	NTID	AAID <u>NT AA</u> score probability
AI7503000987_24258462_c3_982	1529	5301 336 111 200 4.8e-16
phage BK5-T] [SR:Bacteriophage BK5-T ORF'410, 3' end pf cds, 20 ORFs, rep	DNA] [I ressorp:	FN:unidentified] [OR:Lactococcus lactis DB:genpept-phg] [DE:Bacteriophage BK5-T rotein, and Cro repressor protein genes, NT:ORF95; putative] [LE:7452] [RE:7739]
ORF Name	NTID	AAID NT AA score probability
AI7503000987_24275342_c1_718	1530	5302 378 125
Description		
NO-HIT		

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	core	probability	
A17503000987_24320256_f1_134	1531	5303		54			
Description							
NO-HIT							
ORF Name	NTID	AAID	<u>NT</u> Lengthl	<u>AA</u> Length	core	probability	
AI7503000987_24328127_£3_452	1532	5304				6e-10	
Description sp:[LN:YKGC_ECOLI] [AC:P77212] [GN:YKG [SP:P77212] [DB:swissprot] >pir:[LN:He reductase,:ykgC protein] [GN:ykgC] [C >gp:[GI:g1657503] [LN:ECU73857] [AC:U7 [DE:Escherichia coli chromosome minute reductase] [LE:26473] [RE:27825] [DI:C [AC:AE000137:U00096] [PN:putative oxid classified] [OR:Escherichia coli] [DB: section 27 of 400 of the completegenom residues of] [LE:6292] [RE:7644] [DI:C	54756] DR:Esch 73857] es 6-8. complem doreduc genpep ne.] [N	[AC:He erich: [OR:Es of the color of the col	64756] ia coli scheric simila >gp:[GI [GN:ykg 2] [DE:	[PN:pro] [EC:1] hia coli r to S. :g178649 gC] [FN: Escheric	obable .16.1.1 i] [DB: aureus 95] [LA :putati	mercury(II) L] [DB:pir2] genpept-bct1] mercury(II) N:AE000137] ive enzyme; Not	:
ORF Name	NTID	AAID	NT LengthI	AA Length	core	probability	
	533	5305	1386	461 9	45 5.	4e-95]
Description pir:[LN:T00158] [AC:T00158] [PN:amida [EC:3.5] [DB:pir3] >gp:[GI:d103285] [PN:amidase (peptidoglycan hydrolase)] PVL (specific_host:Staphylococcus aure PVL proviral DNA, complete sequence.]	59:g334 OR:beus ATC	1932] acteri] [DB:	LN:AB Lophage genpep	009866] phi PVI t-phg]	[AC:AE L] [SR: DE:Bac	3009866] bacteriophage cteriophage phi	
ORF Name	NTID	AAID	<u>NT</u> LengthI	AA Length	core	probability	
AI7503000987_2438878_£1_126	534	5306	213	70 2	88 2.	3e-25	7
Description	I						J
sp:[LN:ARSC_STAAU] [AC:P30330] [GN:ARS REDUCTASE (ARSENICAL PUMP MODIFIER)] [[AC:D41903] [PN:arsenate reductase,]	[SP:P30	330]	[DB:swi	ssprot]	>pir:[[LN:D41903]	ow

sp:[LN:ARSC_STAAU] [AC:P30330] [GN:ARSC] [OR:STAPHYLOCOCCUS AUREUS] [DE:ARSENATE REDUCTASE (ARSENICAL PUMP MODIFIER)] [SP:P30330] [DB:swissprot] >pir:[LN:D41903] [AC:D41903] [PN:arsenate reductase,] [GN:arsC] [CL:protein-tyrosine-phosphatase, low molecular weight] [OR:Staphylococcus aureus] [EC:1.-.-.] [DB:pir1] >gp:[GI:g150729] [LN:PI2ARSRBC] [AC:M86824] [PN:arsenate reductase] [GN:arsC] [FN:Reduction of arsenate to arsenite] [OR:Plasmid pI258] [SR:Plasmid pI258 DNA] [DB:genpept-bctl] [DE:Plasmid pI258 arsenic resistance operon (arsRBC) genes, completecds.] [LE:1894] [RE:2289] [DI:direct]

ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	score	probability
A17503000987_24401462_c3_921	1535	5307	591	196	367	9.6e-34
Description sp:[LN:YCKF_BACSU] [AC:P42404] [GN:YKD PROTEIN IN TLPC-SRFAA INTERGENIC >pir:[LN:H69760] [AC:H69760] [PN:co [CL:conserved hypothetical protein M >gp:[GI:d1007003:g1438846] [LN:BACYC subtilis] [SR:Bacillus subtilis (str [DB:genpept-bct1] [DE:Bacillus subti chromosomecontaining yckA-H genes.] >gp:[GI:e1182297:g2632631] [LN:BSUBC [OR:Bacillus subtilis] [DB:genpept-bccc] 2 of 21): from 194651 to415810.] [NT [RE:180081] [DI:complement] >gp:[GI:[GN:yckF] [OR:Bacillus subtilis] [SR:[DB:genpept-bcctcccc]] [DB:genpept-bcccccccccccccccccccccccccccccccccccc	REGION onserved MJ1247] CK] [AC:1 cain:168 clis DNA [LE:7448 0002] [Ac oct1] [DI C:similar cd1009614 R:Bacillu	(ORF9) hypot [OR:Ba 030762 trpC2) aroun 3] [RE C:Z991 E:Baci r to h 4:g180 us sub for 2] [SP:Ihetical cillus cillus] [PN:u DNA, cond 28 de :8005] 05:AL00 llus suypothet 5417] tilis cond cond cond cond cond cond cond cond	P42404] I prote subtil unknown clone_l egrees [DI:co 09126] ubtilis cical p [LN:D50 (strain egree r	[DB:sin yeldis] [I GN:ib:lam region mpleme [GN:yeldis] 153] 158 tegion	swissprot] EF] [GN:yckF] DB:pir2] EyckF] [OR:Bacillus Abda DASHI] A of Ent] EkF] [FN:unknown] Lete genome (section AS] [LE:179524] [AC:D50453] ErpC2) DNA] containing
ORF Name A17503000987_24407327_c1_643 Description pir: [LN:E69694] [AC:E69694:S39861:S4 [CL:Escherichia coli ribosomal prote >gp: [GI:e1182036:g2632370] [LN:BSUB0 (BL1)] [GN:rplA] [OR:Bacillus subtil complete genome (section 1 of 21): f [RE:119805] [DI:direct]	ein L1] 0001] [A0 .is] [DB	[OR:Bad C:Z991 :genpe	bosomal cillus 04:AL00 pt-bct1	prote subtil 19126]	in L1] is] [I [PN:ri Bacil]	DB:pir2] .bosomal protein L1 .us subtilis
ORF Name A17503000987_24414050_c2_806 Description gp:[GI:e1296735:g3201550] [LN:SEY171 [OR:Staphylococcus epidermidis] [DB:encoding fibrinogen-bindingprotein,	genpept-	-bct1]	2793 6] [PN: [DE:St	930 fibring	coccus	oinding protein] sepidermidis gene
ORF Name AI7503000987_24415875_c1_660 Description gp:[GI:g1458327] [LN:CELF08F3] [AC:U [SR:Caenorhabditis elegans strain=Br elegans cosmid F08F3.] [LE:1515:1817 [DI:directJoin]	ristol N2	[DB	genper:	OR:Cae	الــــا enorha] [DE:	Caenorhabditis
ORF Name A17503000987_24422175_f2_402 Description NO-HIT	NTID 1539	<u>AAID</u> 5311	NT Length	AA Length	score	probability

ORF Name	NTID	AAID	NT AA LengthLength score probability
AI7503000987_24429643_c2_793	1540	5312	[222] [73] [58] [0.011
Description			
PROTEIN PHAL028] [SP:058584] [DB:swi [PN:hypothetical protein PH0854] [GN [OR:Pyrococcus horikoshii] [DB:pir2] [AC:AP000003:AB009484:AB009485:AB009 hypothetical protein] [GN:PH0854] [O (strain:OT3) DNA] [DB:genpept-bct1]	ssprot] T:PH0854 >gp:[G 486:AB0 PR:Pyroce [DE:Pyro T:simila	>pir:] [CL I:d103 09487: occus ococcu	:hypothetical protein HI0719] 0891:g3257265] [LN:AP000003] AB009488:AB009489] [PN:137aa long horikoshii] [SR:Pyrococcus horikoshii
ORF Name	NTID	AAID	NT AA LengthLength score probability
AI7503000987_24491037_c1_719	1541	5313	495 164
<u>Description</u>			
NO-HIT			
ORF Name	NTID	AAID	NT AA score probability
A17503000987_24500387_c3_961	1542	5314	
Description			
PROTEIN] [SP:P20327] [DB:swissprot] [GN:5.3] [CL:phage T7 gene 2.8 prot [LN:POT3111G] [AC:X17255] [GN:5.3] [>pir:[Ll ein] [Ol OR:Bacte	N:S075 R:phag erioph	
ORF Name	NTID	AAID	NT AA score probability
A17503000987_245953_c3_881	1543	5315	579 192 231 2.5e-19
Description			
<pre>gp:[GI:g1314295] [LN:LMU40604] [AC:U [DB:genpept-bct2] [DE:Listeria monoc [NT:ORF2; putative 19 kDa protein] [</pre>	ytogenes	s ClpC	ATPase (mec) gene, complete cds.]
ORF Name	NTID	AAID	NT AA score probability
A17503000987_24617130_c3_896	1544	5316	3648 1215 5258 0.0
Description			
[GT			

gp:[GI:e187583:g1495791] [LN:SARPOCGEN] [AC:X89233] [PN:DNA-directed RNA polymerase]
[GN:rpoC] [OR:Staphylococcus aureus] [DB:genpept-bct1] [EC:2.7.7.6] [DE:S.aureus DNA
for rpoC gene.] [NT:B' subunit] [SP:P47770] [LE:<1] [RE:>3171] [DI:direct]

ORF Name	NTID	AAID	<u>NT</u> LengthL	AA ength so	core	probability
A17503000987_24648937_c3_924	1545	5317	657 2	218 6	26 3	.4e-61
Description	LJ		<u> </u>	L		
sp:[LN:UNG_BACSU] [AC:P39615] [GN:UN [DE:URACIL-DNA GLYCOSYLASE, (UDG)] [[AC:S39712:H69728] [PN:uracil-DNA g glycosylase] [OR:Bacillus subtilis] [AC:X73124] [GN:ipa-57d] [OR:Bacillu genomic region (325 to 333).] [SP:P3 >gp:[GI:e1186296:g2636332] [LN:BSUBO glycosylase] [GN:ung] [OR:Bacillus s [DE:Bacillus subtilis complete genom [NT:alternate gene name: ipa-57d, yw [DI:complement]	SP:P3961 lycosyla [EC:3.2. s subtil 9615] [I 020] [AC ubtilis] e (secti	L5] [DB Ase, un .2] [Lis] [D LE:5920 C:Z9912 [DB:g ion 20	:swisspg] [GN: DB:pir2 B:genpe 6] [RE: 3:AL009 enpept- of 21):	orot] >pung] ygp: pt-bct1 59883] 126] [P bct1] from 3	pir:[I [CL:ur [GI:g5 L] [DE [DI:d PN:ura [EC:3.	N:S39712] cacil-DNA 680875] [LN:BSGENR] C:B.subtilis direct] ccil-DNA 2.2]
ORF Name	NTID	AAID I	<u>NT</u> LengthL	AA ength so	core	probability
A17503000987_24650332_c3_962	1546	5318	228 7	5		
Description	<u></u>					
NO-HIT						
ORF Name	NTID	AAID I	<u>NT</u> LengthL	AA ength so	core	probability
A17503000987_24650468_f3_522	1547	5319			11 3	.4e-70
Description			L L			
<pre>gp:[GI:d1037645:g4126638] [LN:AB0162 integration and] [OR:bacteriophage pl [DB:genpept-phg] [DE:Bacteriophage pl [LE:25528] [RE:26952] [DI:complement]</pre>	hi-105] hi-105 D	[SR:ba	cteriop	hage ph	ū-105	DNA]
ORF Name	NTID	AAID I	<u>NT</u> LengthLe	AA ength	core	probability
A17503000987_24656552_c2_772	1548	5320	150 4	9 1	49 1	.2e-10
Description			L			
sp:[LN:RL33_THEMA] [AC:P35873] [GN:R:PROTEIN L33] [SP:P35873] [DB:swisspromerous properties of the second properties of t	ot] >gp: otoga ma oteins.] AE001723 [DB:gen	[GI:g4] aritima [SP:P] 3:AE000 apept-bo	07022]] [DB:g: 35873] 512] [P: ct2] [D:	[LN:TMN enpept- [LE:579 N:ribos E:Therm	USGGE bctl]] [RE omal notoga	[] [AC:Z11839] [DE:T.maritima :728] [DI:direct] protein L33] maritima section
ORF Name	NTID	AAID I	<u>NT</u> Length Le	<u>AA</u> ength	ore	probability
A17503000987_24667192_f2_350	1549	5321	183 6	0 2	01 1	.7e-15
Description						
sp:[LN:ARSB_STAAU] [AC:P30329] [GN:AMEMBRANE PROTEIN] [SP:P30329] [DB:sw:[PN:arsenical pump membrane protein] [OR:Staphylococcus aureus] [DB:pirl] [PN:arsenic efflux pump protein] [GN:Plasmid pI258] [SR:Plasmid pI258 resistance operon (arsRBC) genes, con	issprot] [GN:ars >gp:[GI :arsB] [DNA] [D	>pir: B] [Cl :g1507; FN:arse B:genpe	[LN:C41: L:arsen: 28] [LN enic ef: ept-bct:	903] [A ical pu :PI2ARS flux pu 1] [DE:	C:C41 mp men RBC] mp con Plasm	903] mbrane protein] [AC:M86824] mponent (membrane] id pI258 arsenic

ORF Name	NTID	AAID NT AA score probability
AI7503000987_24803462_f1_9	1550	5322 861 286 463 6.4e-44
[DE:HYPOTHETICAL 30.2 KD PROTEIN SLR [AC:S75043] [PN:hypothetical protein PCC 6803] [SR:PCC 6803,] [DB:pir2] [AC:D90910:AB001339] [PN:hypothetical sp. (strain:PCC6803) DNA] [DB:genpep	1717] [n slr17 >gp:[GI l prote t-bct1]	[OR:SYNECHOCYSTIS SP] [SR:PCC 6803,] [SP:P73846] [DB:swissprot] >pir:[LN:S75043] [717] [OR:Synechocystis sp.] [SR:PCC 6803, , [:d1018638:g1652988] [LN:D90910] [cin] [OR:Synechocystis sp.] [SR:Synechocystis [DE:Synechocystis sp. PCC6803 complete [D:slr1717] [LE:37735] [RE:38541] [DI:direct]
ORF Name A17503000987_24823437_f3_521	<u>NTID</u>	AAID NT AA score probability S323 495 164 71 0.044
[DE:Staphylococcus plasmid pC221.] [in] [SP:P03866] [LE:4287] [RE:>4555] [AC:X02529] [OR:Staphylococcus aureu	NT:pot. [DI:di: s] [DB:q] [NT:po	R:Staphylococcus aureus] [DB:genpept-bct1] reading-frame C (aa 1-90) (4555 is 2nd base rect] >gp:[GI:e190060:g1333818] [LN:SAPC221] genpept-bct1] [DE:Staphylococcus aureus oct. orfB (aa 1-92) (4557 is 2nd base in DI:direct]
ORF Name	NTID	AAID NT AA score probability
AI7503000987_24854637_c2_804	1552	5324 903 300 404 1.1e-37
Description		
[CL:hypothetical protein ywpJ] [OR:B >gp:[GI:e1184491:g2636110] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-b 19 of 21): from 3597091to 3809700.] [RE:98190] [DI:direct] >gp:[GI:e3080 [OR:Bacillus subtilis] [DB:genpept-b genes.] [NT:product similar to Bacil [DI:complement] >gp:[GI:e1184491:g26 [FN:unknown] [OR:Bacillus subtilis]	acillus 019] [A0 ct1] [DI [NT:sim: 93:g1894 ct1] [DI lus subt 36110] [DB:genp	hypothetical protein ywtE] [GN:ywtE] subtilis] [DB:pir2] C:Z99122:AL009126] [GN:ywtE] [FN:unknown] E:Bacillus subtilis complete genome (section dilar to hypothetical proteins] [LE:97330] 4770] [LN:BSZ92954] [AC:Z92954] [GN:ywtE] E:B.subtilis yws[A,B,C,D,E,F,G] and gerBC ctilis YxeH and YcsE] [LE:4292] [RE:5152] [LN:BSUB0019] [AC:Z99122:AL009126] [GN:ywtE] pept] [DE:Bacillus subtilis complete genome] [NT:similar to hypothetical proteins]
ORF Name	NTID	AAID NT AA score probability
A17503000987_24855325_c3_936	1553	5325 129 42
Description NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000987_25398262_f3_622 Description NO-HIT	1554	5326 132 43
ORF Name	NTID	AAID NT AA score probability
A17503000987_25478801_£3_446	1555	5327 [147] [48]
Description		
NO-HIT		

AANTAAID ORF Name NTID score probability LengthLength A17503000987 25509692 c3 972 1556 5328 498 165 .4e-11 Description gp:[GI:e244846:g2764870] [LN:BSPP1GENM] [AC:X97918] [OR:Bacteriophage SPP1] [DB:genpept-phq] [DE:Bacteriophage SPP1 complete nucleotide sequence.] [NT:gene 17.5] [LE:11342] [RE:11881] [DI:direct] NT AA AAID score ORF Name NTID probability LengthLength A17503000987_25578827_c3_897 5329 261 86 3.0e-30 1557 334 Description sp:[LN:YBXF STAAU] [AC:Q53602] [OR:STAPHYLOCOCCUS AUREUS] [DE:PROBABLE RIBOSOMAL PROTEIN IN RPSL 5'REGION (FRAGMENT)] [SP:Q53602] [DB:swissprot] >gp:[GI:g706920] [LN:SAU20869] [AC:U20869] [PN:unknown] [FN:unknown] [OR:Staphylococcus aureus] [DB:genpept-bct2] [DE:Staphylococcus aureus ribosomal protein S12 (rpsL) gene, completecds, ribosomal protein S7 (rpsG) and ORF 1 genes, partial cds.] [NT:ORF 1] [LE:<1] [RE:320] [DI:direct] NT AΑ <u>sco</u>re ORF Name NTID AAID probability LengthLength AI7503000987_25579662_£3_575 1558 1221 8.0e-85 Description pir: [LN:H69817] [AC:H69817] [PN:aminoacylase homolog yhaA] [GN:yhaA] [CL:hippurate hydrolase] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1183009:g2633343] [LN:BSUB0006] [AC:Z99109:AL009126] [GN:yhaA] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 6 of 21): from 999501 to1209940.] [NT:similar to aminoacylase] [LE:80123] [RE:81313] [DI:complement] NT ORF Name NTID AAID <u>sco</u>re probability LengthLength A17503000987 25660937 c3 990 498 1559 5331 165 3.5e-13 Description sp:[LN:OPUC BACSU] [AC:P46922] [GN:OPUAC] [OR:BACILLUS SUBTILIS] [DE:GLYCINE BETAINE-BINDING PROTEIN PRECURSOR] [SP:P46922] [DB:swissprot] >pir:[LN:I40537] [AC:I40537:F69669] [PN:qlycine betaine ABC transporter (glycine betaine-binding protein) opuAC precursor] [GN:opuAC] [OR:Bacillus subtilis] [DB:pir2] >qp:[GI:e1182252:q2632586] [LN:BSUB0002] [AC:Z99105:AL009126] [PN:qlycine betaine ABC transporter (glycine) [GN:opuAC] [FN:glycine betaine transport (osmoprotection)] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 2 of 21): from 194651 to415810.] [SP:P46922] [LE:128023] [RE:128904] [DI:direct] >gp:[GI:d1009569:g1805372] [LN:D50453] [AC:D50453] [PN:glycine betain-binding protein precursor] [GN:opuAC] [OR:Bacillus subtilis] [SR:Bacillus subtilis (strain:168 trpC2) DNA] [DB:genpept-bct1] [DE:Bacillus subtilis DNA for 25-36 degree region containing theamyE-srfA region, complete cds.] [LE:2326] [RE:3207] [DI:direct] >gp:[GI:g984805] [LN:BSU17292] [AC:U17292] [PN:glycine betaine-binding protein precursor] [GN:opuAC]

[OR:Bacillus subtilis] [DB:genpept-bct2] [DE:Bacillus subtilis ATPase (opuAA), transmembrane protein (opuAB) andglycine betaine-binding protein precursor (opuAC)

genes, completecds.] [LE:2332] [RE:3213] [DI:direct]

ORF Name	NTID	AAID NT AA score probability
A17503000987_25667753_c2_773	1560	
ANTITERMINATION PROTEIN NUSG] [SP:P: [PN:transcription antitermination fantitermination factor nusG] [OR:State [LN:SCSECE] [AC:X76134] [GN:nusG] [OR:NusG] [CO:X76134]	36264] [actor nu aphyloco DR:Staph	OR:STAPHYLOCOCCUS CARNOSUS] [DE:TRANSCRIPTION [DB:swissprot] >pir:[LN:S38870] [AC:S38870] usG] [GN:nusG] [CL:transcription occus carnosus] [DB:pir2] >gp:[GI:g426473] hylococcus carnosus] [DB:genpept-bct1] [SP:P36264] [LE:331] [RE:879] [DI:direct]
ORF Name	NTID	AAID NT AA score probability
A17503000987_25972086_f1_108 Description NO-HIT	1561	5333 135 44
ORF Name	NTID	$\frac{\text{AAID}}{\text{LengthLength}} \frac{\text{NT}}{\text{score}} \text{probability}$
Description gp:[GI:e244712:g2764863] [LN:BSPP1GI [DB:genpept-phg] [DE:Bacteriophage S [LE:9322] [RE:9651] [DI:direct]		[5334] [363] [120] [103] [9.0e-06] C:X97918] [OR:Bacteriophage SPP1] mplete nucleotide sequence.] [NT:gene 16]
ORF Name	NTID	AAID NT AA score probability
AI7503000987_26206687_c1_642	1563	5335 210 69 278 2.6e-24
TRANSLOCASE SECE SUBUNIT] [SP:006442 [AC:U96619] [PN:SecE] [GN:secE] [OR	2] [DB:s :Staphyl :SecE (s	secE), NusG (nusG) and RplK(rplK) genes,
ORF Name	NTID	AAID NT AA score probability
A17503000987_26290912_f3_547	1564	5336 882 293 665 2.5e-65
[OR:Bacillus subtilis] [DB:pir2] >gr [AC:Z99105:AL009126] [GN:yciA] [FN:u [DE:Bacillus subtilis complete genor [NT:similar to hypothetical proteins >gp:[GI:d1009603:g1805406] [LN:D5049 [SR:Bacillus subtilis (strain:168 to	p:[GI:e1 unknown] ne (sect s] [LE:1 53] [AC: cpC2) DN] [OR:Bacillus subtilis] [DB:genpept-bct1] tion 2 of 21): from 194651 to415810.]
ORF Name	NTID	AAID NT AA score probability
AI7503000987_26306568_c1_724	1565	
		4] [PN:pXO1-90] [OR:Bacillus anthracis] irulence plasmid PX01, complete sequence.]

ORF Name	NTID	AAID	NT AA LengthLength score probability
AI7503000987_26350125_c2_834	1566	5338	79 0.041
Description		L	
pir:[LN:C69161] [AC:C69161] [PN:sen [OR:Methanobacterium thermoautotroph [AC:AE000831:AE000666] [PN:sensory t [OR:Methanobacterium thermoautotroph thermoautotrophicum from bases 40481 genome.] [NT:Function Code:12.12 - C. [DI:complement]	icum] [ransduc icum] [7 to 41	DB:pir: tion h: DB:gen; 5582(se	2] >gp:[GI:g2621537] [LN:AE000831] istidine kinase] [GN:MTH468] pept-bctl] [DE:Methanobacterium ection 37 of 148) of the complete
ORF Name	NTID	AAID	NT AA LengthLength
AI7503000987_26360327_c1_725	1567	5339	552 183 515 2.0e-49
	:[GI:d1 VL] [SR ATC] [032887 :bacte: DB:gen;	pept-phg] [DE:Bacteriophage phi PVL
ORF Name	NTID	AAID	NT <u>AA</u> LengthLength
AI7503000987_26369082_c2_837	1568	5340	252 83
Description			
NO-HIT	or an extra contract to the co		
ORF Name	NTID	AAID	NT AA LengthLength probability
AI7503000987_26375952_c1_714	1569	5341	627 208 223 1.7e-18
Description gp:[GI:e1254413:g2924242] [LN:BTP901- [OR:Bacteriophage TP901-1] [DB:genpe] [NT:ORF11] [LE:5545] [RE:6168] [DI:d	pt-phg]		
ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000987_26569377_c2_810	1570	5342	639 212 625 4.4e-61
Description			
sp:[LN:HUMS_BACSU] [AC:P42405:O31477] [DE:3-HEXULOSE 6-PHOSPHATE FORMALDEHT >pir:[LN:A69761] [AC:A69761] [PN:D-a yckG] [GN:yckG] [OR:Bacillus subtil: [LN:BSUB0002] [AC:Z99105:AL009126] [OB:genpept-bct1] [DE:Bacillus subtil: to415810.] [NT:similar to D-arabino : [RE:180719] [DI:complement]	YDE LYA arabino is] [DB GN:yckG lis com	SE)] [S 3-hexu :pir2]] [FN:u plete g	SP:P42405:O31477] [DB:swissprot] Alose 6-phosphate formaldeh homolog >gp:[GI:e1182298:g2632632] Anknown] [OR:Bacillus subtilis] genome (section 2 of 21): from 194651
ORF Name	NTID	AAID	NT AA LengthLength score probability
AI7503000987_26734625_f1_89	1571	5343	351 116 210 4.1e-17
Description		<u> </u>	
<pre>gp:[GI:e308969:g2292761] [LN:BSFI21LY [OR:Streptococcus thermophilus bacter thermophilus bacteriophage Sfi21 DNA; [RE:6076] [DI:complement]</pre>	riophag	e Sfi21	[PN:cI-like repressor] [GN:orf127] [DB:genpept-phg] [DE:Streptococcus le, 8141 bp.] [NT:putative] [LE:5693]

ORF Name	NTID	AAID	NT AA score probability
AI7503000987_26772801_£1_91	1572	5344	504 167 168 6.1e-12
Description pir: [LN:A71608] [AC:A71608] [PN:problem: [GN:PFB0710c] [OR:Plasmodium falcipated [AC:AE001413:AE001362] [PN:predicted [OR:Plasmodium falciparum] [SR:malar: [DE:Plasmodium falciparum chromosome [NT:predicted by GlimmerM] [LE:4351]	arum] [integr ia para 2, sec	DB:pir: al mem site P tion 50	2] >gp:[GI:g3845257] [LN:AE001413] brane protein] [GN:PFB0710c] . falciparum] [DB:genpept-inv2] 0 of 73 of thecomplete sequence.]
ORF Name	NTID	AAID	NT AA score probability
A17503000987_26839638_c1_750	1573	5345	1314 437 1429 2.8e-146
Description			
sp:[LN:YWFO_BACSU] [AC:P39651] [GN:YWFO_BECSU] [AC:P39651] [GN:YWFO_DE:HYPOTHETICAL 51.0 KD PROTEIN IN FOUR PR	PTA 3'R [PN:yw 3:pir2] nknown] e (sect milar t L567] [genpep	EGION] fO prot >gp:[0 [OR:Ba ion 20 o] [SP: LN:BSUV t-bct1]	[SP:P39651] [DB:swissprot] tein:hypothetical protein ipa-93d] GI:e1186260:g2636296] [LN:BSUB0020] acillus subtilis] [DB:genpept-bct1] of 21): from 3798401to 4010550.] :P39651] [LE:59934] [RE:61235] WFO] [AC:Z80355] [PN:Unknown] [DE:B.subtilis ywfO, ywgA and ywgB
ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000987_26854757_c3_886	1574	5346	
sp:[LN:CYSE_STAXY] [AC:P77985] [GN:CY [DE:SERINE ACETYLTRANSFERASE, (SAT)] >gp:[GI:e261410:g1514656] [LN:SXCYSER [GN:cysE] [OR:Staphylococcus xylosus] cysE gene, genomic region.] [SP:P7798	SP:P7 REG] [A DB:go	7985] C:Y0761 enpept-	[DB:swissprot] L4] [PN:serine O-acetyltransferase] -bctl] [EC:2.3.1.30] [DE:S.xylosus
ORF Name	NTID	AAID	NT AA LengthLength score probability
AI7503000987_27318_c2_877	1575	5347	1198 8.3e-122
	[SP:P: ol dehyd drogena [LN] [LN genpept	20368] drogena ase hom :ZMOADH -bctl]	[DB:swissprot] >pir:[LN:A35260] ase, 1] [GN:adhA] [CL:alcohol mology] [OR:Zymomonas mobilis] HA] [AC:M32100] [OR:Zymomonas mobilis] [DE:Z.mobilis alcohol dehydrogenase I
ORF Name	NTID	AAID	NT AA LengthLength score probability
AI7503000987_276590_c2_821	1576	5348	375 124 279 2.0e-24
Description pir:[LN:H70070] [AC:H70070] [PN:hypound	g263629 ubtilis	97] [LN s] [DB:	J:BSUB0020] [AC:Z99123:AL009126] genpept-bct1] [DE:Bacillus subtilis

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
AI7503000987_2847887_£2_351	1577	5349	576	191		2.6e-79
Description		L				
sp:[LN:ARSB_STAAU] [AC:P30329] [GN:A MEMBRANE PROTEIN] [SP:P30329] [DB:sw [PN:arsenical pump membrane protein] [OR:Staphylococcus aureus] [DB:pir1] [PN:arsenic efflux pump protein] [GN [OR:Plasmid pI258] [SR:Plasmid pI258 resistance operon (arsRBC) genes, co	issprot [GN:ar >gp:[G :arsB] DNA] [1] >pir sB] [I:g150 [FN:ar DB:gen	:[LN:C4 CL:arse 728] [I senic e pept-bo	11903] enical LN:PI2A efflux ct1] [D	[AC:C4 pump n RSRBC] pump c E:Plas	nembrane protein] [AC:M86824] component (membrane) smid pI258 arsenic
ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	score	probability
A17503000987_29320127_f3_442	1578	5350	135	44		
Description NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000987_29694425_f1_142	1579	5351	852	283	418	3.8e-39
Description gp:[GI:g1502421] [LN:BSU59433] [AC:U						
PlsX (plsX), malonyl-CoA:Acyl carrie carrier protein reductase(fabG) gene gene,partial cds.] [NT:also called 3 [DI:direct]	s, comp	lete c	ds, and	l acyl	carrie	r protein (acpP)
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000987_30178137_c1_706	1580	5352		43		
Description NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000987_30267937_£3_532	1581	5353	123	40		
Description NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000987_30742332_c2_841	1582	5354	228	75	76	0.012
Description pir: [LN:D71169] [AC:D71169] [PN:hype [OR:Pyrococcus horikoshii] [DB:pir2] [AC:AP000002:AB009475:AB009476:AB0094 hypothetical protein] [GN:PH0552] [OX (strain:OT3) DNA] [DB:genpept-bct1] 287001-544000 nt. position(2/7).] [NY [LE:205880] [RE:206371] [DI:direct]	>gp:[G] 477:AB00 R:Pyroco [DE:Pyro	:d1036)9478:) occus l	0584:g3 AB00947 horikos s horik	256958; 9:AB00; hii] [: oshii (] [LN: 9480] SR:Pyr OT3 ge	AP000002] [PN:163aa long ococcus horikoshii nomic DNA,

ORF Name	NTID	AAID	NT Length Le	AA ength	re p	robabili	.ty
AI7503000987_31256916_c2_774	1583	5355	[507] [I	68 68:	5.1	e-67	
Description			J L L.				
pir:[LN:S38871] [AC:S38871] [PN:rik ribosomal protein L11] [OR:Staphyloo [LN:SCSECE] [AC:X76134] [PN:L11 prot [DB:genpept-bct1] [DE:S.carnosus sec [RE:1483] [DI:direct]	coccus ca cein] [Gl	arnosus N:rplK]	s] [DB:p:] [OR:Sta	ir2] >gp aphyloco	:[GI:g	3581638] carnosus]	
ORF Name	NTID	AAID	<u>NT</u> LengthLe	AA ength	re p	robabili	.ty
A17503000987_31334838_c3_993	1584	5356	519 1	72 544	1.7	e-52	
Description			J L			 	
pir:[LN:E70057] [AC:E70057] [PN:hypsubtilis] [DB:pir2] >gp:[GI:e267625:[GN:ywhD] [OR:Bacillus subtilis] [DEchromosomal region.] [LE:3358] [RE:3 [LN:BSUB0020] [AC:Z99123:AL009126] [DB:genpept-bct1] [DE:Bacillus subti3798401to 4010550.] [LE:52838] [RE:5	g1565239 3:genpept 8876] [D: [GN:ywhD] .lis comp	9] [LN: t-bct1] I:dired] [FN:u plete g	:BSTHRZ] [DE:B.; ct] >gp: unknown] genome (s	AC:Z80 subtilis [GI:e118] [OR:Bac	360] thrZ 6252:g illus	[PN:Unkno downstre j2636288] subtilis	own] eam s]
ORF Name	NTID	AAID	<u>NT</u> LengthLe	AA ngth sco	re p	robabili	ty
AI7503000987_3134386_f3_616	1585	5357	228 7				
Description							
NO-HIT							
ORF Name	NTID	AAID	<u>NT</u> LengthLe	AA ength sco	re p	robabili	ty
A17503000987_31353377_c2_785	1586	5358	474 1	57 703	2.4	e-69	
Description			·				
<pre>gp:[GI:d1039003:g4512400] [LN:AB0175 halodurans] [SR:Bacillus halodurans halodurans C-125 genomic DNA, 32 kb (identity of 84% to B. subtilis)] [</pre>	(strain: fragment	:C-125) :, comp	DNA] [I	DB:genpe .] [NT:r]	pt-bct psG ho	1] [DE:E	Bacillus
ORF Name	NTID	AAID	<u>NT</u> LengthLe	AA ength	re p	robabili	ty
A17503000987_31562_f2_326	1587	5359	123 40	רס			
Description							
NO-HIT							
ORF Name	NTID	AAID	<u>NT</u> LengthLe	AA ngth sco	re p	robabili	ty
AI7503000987_32031437_c1_736	1588	5360	2706 90		3 4.3	e-118	1
Description	L		J				
sp:[LN:VG12_BPPZA] [AC:P07537] [GN:1 PROTEIN (LATE PROTEIN GP12)] [SP:P07 [PN:gene 12 protein] [GN:12] [CL:ph >gp:[GI:g216061] [LN:PZACG] [AC:M118 [OR:Bacteriophage PZA] [SR:Bacteriop PZA (from B.subtilis), complete geno	537] [DE age PZA 13:M1390 hage PZA	3:swiss gene 1 04:M139 A DNA]	sprot] >p .2 protei .05] [PN: 	oir:[LN:Vin] [OR:p pre-neck pept-phg]	MBP12 phage appe [DE:] [AC:G2 PZA] [DB ndage pr Bacterio	4831] :pir1] otein]

ORF Name	NTID	AAID LengthLength score probability
AI7503000987_32251_£1_98	1589	5361 129 42
Description		
NO-HIT		
ORF Name	NTID	AAID NT AA core probability
A17503000987_32315907_f3_595	1590	5362 126 41
Description		
NO-HIT		
ORF Name	NTID	AAID <u>NT AA</u> score probability
AI7503000987_32595152_c1_739	1591	5363 402 133
Description		
NO-HIT	er in part of the state of the	
ORF Name	NTID	AAID NT AA score probability
AI7503000987_3306563_c1_688	1592	5364 366 121
<u>Description</u>		
NO-HIT	- 00 D	
ORF Name	NTID	AAID NT AA score probability
AI7503000987_33235050_f2_235 Description	1593	5365 507 168 92 0.0014
pir: [LN:D70063] [AC:D70063] [PN:hypsubtilis] [DB:pir2] >gp: [GI:e1184569 [GN:ywnA] [FN:unknown] [OR:Bacillus complete genome (section 19 of 21): [DI:complement] >gp: [GI:e269486:g159 [GN:ywnA] [OR:Bacillus subtilis] [DB downstream DNA.] [LE:2688] [RE:3089] [LN:BSUB0019] [AC:Z99122:AL009126] [:g263618 subtilis from 359 2697] [3 :genpept [DI:din GN:ywnA] omplete	t-bct1] [DE:B.subtilis urease operon and rect] >gp:[GI:e1184569:g2636188]] [FN:unknown] [OR:Bacillus subtilis] genome (section 19 of 21): from 3597091to
ORF Name	NTID	AAID NT AA score probability
AI7503000987_33394062_f1_179	1594	5366 129 42
Description		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000987_33414693_c2_820	1595	5367 1089 362 150 6.9e-16
[DE:MEVALONATE KINASE, (MK)] [SP:Q58 [PN:mevalonate kinase,] [OR:Methanoc [MP:FOR1026197-1027135] >gp:[GI:g15 [PN:mevalonate kinase] [GN:MJ1087] [487] [DI occus ja 91731] OR:Metha 93 of 1	[LN:U67551] [AC:U67551:L77117] anococcus jannaschii] [DB:genpept-bct2] L50 of the complete genome.] [NT:similar to

ORF Name	NTID	AAID <u>NT AA</u> LengthLength score probability
A17503000987_33600035_c1_653	1596	5368 2118 705 3427 0.0
Description		
[GN:fus] [FN:translation elongation	factor]	C:AJ237696] [PN:elongation factor G (EF-G)] [OR:Staphylococcus aureus] fus gene.] [LE:83] [RE:2164] [DI:direct]
ORF Name	NTID	AAID NT AA score probability
A17503000987_33756503_c2_830	1597	5369 273 90 265 6.2e-23
	re A2] [I	Y12813] [PN:hypothetical protein] [GN:orfB] DB:genpept-phg] [DE:Bacteriophage A2 rep, complement]
ORF Name	NTID	AAID NT AA score probability
A17503000987_33786251_c2_865	1598	5370 246 81
Description		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000987_33869193_c1_726	1599	5371 183 60 130 1.2e-08
Description sp:[LN:RINB_BPPHA] [AC:Q03183] [GN:F	INB] [OI	P.RACTEDIODHAGE DHI-11] [DE.TDANSCRIDTIONAL
activator RinB] [OR:phage phi-11] [I [PN:Bacteriophage phi-11 int gene ac	sprot] : B:pir2] tivator] bacteric	<pre>>pir:[LN:A49703] [AC:A49703] [PN:int gene >gp:[GI:g166161] [LN:BPHRINAB] [AC:L07580]] [GN:rinB] [FN:Activate bacteriophage ophage phi 11] [SR:Bacteriophage phi-11 DNA] inA and rin B genes, required for</pre>

ORF Name	NTID	AAID	NT Lengtl	<u>AA</u> 1Length	score	probabili	ty
A17503000987_34242202_c3_956	1601	5373	270	89	78	0.017	
Description Sp:[LN:VP8_VARV] [AC:P33039] [GN:L4R PRECURSOR (25 KD MAJOR CORE PROTEIN) pir:[LN:A36845] [AC:A36845:S33090] [CL:vaccinia virus 28K protein] [OR: [LN:S55844] [AC:S55844] [PN:glycopro [SR:Variola major virus India-1967] virus, India-1967, Genomic, 19 genes [LE:3187] [RE:3942] [DI:direct] >gp: major virus] [SR:Variola major virus [DE:Variola major virus (strain Bang vaccinia virus CDS L4R (structural) [LN:VVGGAA] [AC:X6919] [GN:M4R] [OR DNA complete genome.] [SP:P33039] [L [LN:VVHINDQKH] [AC:X67119] [PN:struc [DB:genpept-vrl] [DE:Variola virus ([SP:P33039] [LE:19835] [RE:20590] [D	::M4R] (P25K) [PN:M4 variola tein VI [DB:ger ,18029 [GI:g43 (strai ladesh- [LE:705 ::Variol E:69950 tural r HindIII	[OR:VAR] [SP: 4R prote a virus P8] [GN npept-v nt].] 38994] in Bang -1975) 6 583] [RI la virus	IOLA VP33039 ein:st [DB:::L4R] rl] [D [NT:Th [LN:VA ladesh comple E:7133 s] [DB 70705] VP8]	IRUS] [] [DB:s ructura pir2] > [OR:Var E:G9R is sequ RCG] [A -1975) te genc 8] [DI: :genpep [DI:di [GN:L4R	[DE:ST] Swissp: al properties of the properties	RUCTURAL PRO rot] tein VP8] I:g262435] major virus] [variola maj comes from F 579] [OR:Var [DB:genpept- [NT:homolog t] >gp:[GI:g62 [DE:Variola OR:Variola	or ig. 2.] iola vrl] of 297256] a virus 352] virus]
ORF Name AI7503000987_34242202_f1_156 Description sp:[LN:Y076_MYCPN] [AC:P75555] [OR:M MG076 HOMOLOG] [SP:P75555] [DB:swiss homolog G07_orf138] [OR:Mycoplasma p 29342,] [DB:pir2] >gp:[GI:g1674320] [GN:G07_orf138] [OR:Mycoplasma pneum section 59 of 63 of the complete gen [LE:168] [RE:584] [DI:direct]	prot] > neumoni [LN:ME oniae]	pir:[L] lae] [S] PAE0000! [DB:ge	JMONIA N:S739 R:ATCC 59] [Adapept-]	43] [AC 29342, C:AE000 bct2] [HYPOTH ::S7394 , ATC	13] [PN:MG0 CC 29342] [S 00089] coplasma pne	EIN 76 R:ATCC umoniae
ORF Name	NTID			<u>AA</u> 1Length	score	probabili	<u> </u>
A17503000987_34407053_f3_433	1603	5375	783	260	564	1.3e-54	
Description pir:[LN:S76797] [AC:S76797] [PN:hyp protein MJ0165: phosphoribosylaminoi [OR:Synechocystis sp.] [SR:PCC 6803, pgp:[GI:d1019442:g1653798] [LN:D9091 [OR:Synechocystis sp.] [SR:Synechocy [DE:Synechocystis sp. PCC6803 comple [NT:ORF_ID:s111489] [LE:107538] [RE:	midazol , PCC 6] [AC: stis sp te geno	le carbo 6803] D90916 D. (stra Dome, 26,	oxylase [SR:PC0 :AB0011 ain:PC0 '27, 32	e catal C 6803, 339] [P C6803) 270710-	ytic o] [DE N:hypo DNA]	chain homolog 3:pir2] othetical pro [DB:genpept-]	gy] otein]
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probabilit	<u> </u>
A17503000987_34609703_c3_963	1604	5376	198	65	110	1.6e-06	
Description							
pir:[LN:T00185] [AC:T00185] [PN:hypophage phi PVL] [DB:pir3] >gp:[GI:d10]		-					ceus

[OR:bacteriophage phi PVL] [SR:bacteriophage phi PVL (specific_host:Staphylococcus aureus ATC] [DB:genpept-phg] [DE:Bacteriophage phi PVL proviral DNA, complete sequence.] [NT:orf 52] [LE:37356] [RE:37604] [DI:direct]

ORF Name	NTID	AAID	NT Length I	Length score	probability	
AI7503000987_34651555_c1_734	1605	5377	3123	1040 267	2.8e-24	
Description			. ســـــــا د			
gp:[GI:e244718:g2764872] [LN:BSPP1GEN						
[DB:genpept-phg] [DE:Bacteriophage SP: [LE:12264] [RE:13373] [DI:direct]	br comb	tere	nucleot.	ide sequence	e.] [NT:gene 18]	
		Marie of the second of				
	NTID	AAID	NT Length I	AA Length	probability	
AI7503000987_35162800_c3_885	1606	5378	1506	501 1701	4.2e-175	
Description						
sp:[LN:SYE_BACSU] [AC:P22250] [GN:GLT:						
[DE:(GLURS)] [SP:P22250] [DB:swissprof [PN:glutamatetRNA ligase, gltX:glutamatetrnA ligase, gltX:glutamate						
[CL:glutamatetRNA ligase:glutamine-						
[EC:6.1.1.17] [DB:pirl] >gp:[GI:d1005						
[PN:glutamyl-tRNA synthetase] [GN:glt]						
(sub_species:Marburg, strain:168) DNA						
kilobase region of replication origin >gp:[GI:g143006] [LN:BACGLTXA] [AC:M5						
[GN:gltX] [OR:Bacillus subtilis] [SR:1						
[EC:6.1.1.17] [DE:Bacillus subtilis g						
[LE:1] [RE:1452] [DI:direct] >gp:[GI:						
[PN:glutamyl-tRNA synthetase] [GN:glt]						
<pre>(strain 168T) DNA] [DB:genpept-bct1] (gltX), serineacetyltransferase (cysE)</pre>						
complete cds's.] [LE:1706] [RE:3157]						
[LN:BSUB0001] [AC:Z99104:AL009126] [PI						3
subtilis] [DB:genpept-bct1] [EC:6.1.1	.17] [D	E:Bac	illus sı	ubtilis com	plete genome	
(section 1 of 21): from 1 to213080.]					=	
>gp:[GI:g2653659] [LN:U49789] [AC:U49° subtilis] [DB:genpept-bct2] [DE:Bacil]						
complete cds.] [LE:1] [RE:1452] [DI:d:		CIIIS	grucam	YI-CKNA SYIN	metase gene,	
		·				_
ORF Name	NTID	AAID	NT	AA score	probability	
A17503000987_35350062_£2_317	1607	5379	LengthI	46		
Description	النت		ا لــــــال			
NO-HIT						
			11.11.234		The second of th	_
ORF Name	NTID	AAID	NT LengthL	AA score	probability	
AI7503000987_35445875_c1_728	1608	5380			0.016	
Description			JL			
pir:[LN:C70306] [AC:C70306] [PN:conse	erved h	ypothe	etical r	orotein ag ()64c] [GN:ag 064c]	
[CL:conserved hypothetical secreted pr						

pir:[LN:C70306] [AC:C70306] [PN:conserved hypothetical protein aq_064c] [GN:aq_064c] [CL:conserved hypothetical secreted protein HP0320] [OR:Aquifex aeolicus] [DB:pir2] >gp:[GI:g2982828] [LN:AE000672] [AC:AE000672:AE000657] [PN:hypothetical protein] [GN:aq_064c] [OR:Aquifex aeolicus] [DB:genpept-bct2] [DE:Aquifex aeolicus section 4 of 109 of the complete genome.] [LE:2190] [RE:2423] [DI:direct]

ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	score	probability
A17503000987_36054813_c2_833	1609	5381		128	79	0.034
Description pir: [LN:S41186] [AC:S43812:S41186] >gp: [GI:e244746:g2764918] [LN:BSPP1G [DB:genpept-phg] [DE:Bacteriophage S [LE:36342] [RE:36722] [DI:direct] >g [OR:Bacteriophage SPP1] [DB:genpept-for products required forreplication	ENM] [A0 PP1 comp p:[GI:g4 phg] [DI	C:X979 plete 139643 E:B.su	18] [OR nucleot] [LN:B btilis	:Bacte ide se SSPP1] phage	riopha quence [AC:X	age SPP1] e.] [NT:gene 39] K67865] [GN:39] DNA sequence coding
ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	score	probability
A17503000987_36134715_c3_887	1610	5382		256	666	2.0e-65
Description		L	J L		LJ	
sp:[LN:YACO_BACSU] [AC:Q06753] [GN:Y. [DE:HYPOTHETICAL TRNA/RRNA METHYLTRA: >pir:[LN:S66124] [AC:S66124:G69741:I. [GN:yacO] [CL:conserved hypothetica >gp:[GI:d1005871:g467483] [LN:BAC180] [SR:Bacillus subtilis (sub_species:M. subtilis DNA, 180 kilobase region of [DI:direct] >gp:[GI:e1182029:g263236 [FN:unknown] [OR:Bacillus subtilis] genome (section 1 of 21): from 1 to2 [SP:Q06753] [LE:115266] [RE:116015]	NSFERASE 40000] l protei K] [AC:I arburg, replica 3] [LN:E [DB:genr 13080.]	E YACO [PN:con HIOs 226185] strain ation of BSUB000 pept-be [NT:s:	,] [SP: conserve 860] [O] [PN:u n:168) crigin. col] [AC ctl] [D	Q06753 d hypo R:Baci nknown DNA] [] [LE: :Z9910 E:Baci	DB: thetic llus s [OR: DB:ger 178868 4:AL00 llus s	swissprot] cal protein yac0] subtilis] [DB:pir2] Bacillus subtilis] upept-bct1] [DE:B. B] [RE:179617] 09126] [GN:yac0] subtilis complete
ORF Name	NTID	AAID	<u>NT</u> Length:	<u>AA</u> Length	score	probability
A17503000987_36228563_c3_960	1611	5383		232	110	0.00063
Description gp:[GI:g2865254] [LN:AF008237] [AC:A: [OR:Mitochondrion Chlamydomonas eugai [DB:genpept-pln2] [DE:Chlamydomonas eugai [NT:orf306; intronic ORF; formerly i.	metos] (eugameto	[SR:Ch]	lamydom ochondr	onas e ion, c	ugamet omplet	os] e genome.]
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000987_36521067_£3_540	1612	5384	204	67		
Description NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Lengthl	<u>AA</u> Length	score	probability
A17503000987_3908462_c1_703	1613	5385		47		
Description NO-HIT		k				
ORF Name	NTID	AAID	<u>NT</u> Lengthl	AA Length	score	probability
AI7503000987_3909643_t2_217	1614	5386		398	719	4.8e-71
Description	7005 775				-	
gp:[GI:g4981502] [LN:AE001759] [AC:AI protein] [GN:TM0964] [OR:Thermotoga r section 71 of 136 of the complete ger 60.38;] [LE:5733] [RE:6941] [DI:direction of the complete ger for the complete ger f	maritima nome.] [[DB:	genpep	t-bct2] [DE:	Thermotoga maritima

ORF Name	NTID	AAID	<u>NT</u> Lengthl	Length score	probability
AI7503000987_3912503_c1_698	1615	5387	1500	499 331	1.5e-28
Description		L	JII	—————	
pir:[LN:E69825] [AC:E69825] [PN:ami	ino acid	trans	porter 1	homolog vhd	G] [GN:vhdG]
[CL:arginine permease] [OR:Bacillus					
[LN:BSUB0006] [AC:Z99109:AL009126]			_		-
[DB:genpept-bct1] [DE:Bacillus subti	llis com	plete	genome	(section 6 d	of 21): from 999501
to1209940.] [NT:similar to amino aci		_			
>gp:[GI:e325026:g2226202] [LN:BSY140					-
[OR:Bacillus subtilis] [DB:genpept-k					
72 to 75 degrees: spoVRto sspB.] [NT [LE:9978] [RE:11375] [DI:direct]	::Simila	rity t	o numan	retroviral	receptor (PIR)
[LE:9976] [RE:II375] [DI:direct]					4114
ORF Name	NTID	AAID	NT	AA score	probability
	11111		LengthI	Length Door	probability
A17503000987_3945333_c2_782	1616	5388	168	55	
Description					
NO-HIT					
			NT	AA	
ORF Name	NTID	AAID	LengthI	ength score	probability
A17503000987_3961590_c3_922	1617	5389			9.5e-66
Description					
pir:[LN:H69831] [AC:H69831] [PN:ace	tyl-CoA	C-ace	tyltrans	sferase homo	olog yhfS] [GN:yhfS
] [OR:Bacillus subtilis] [DB:pir2] >					
[AC:Z99109:AL009126] [GN:yhfs] [FN:u					
[DE:Bacillus subtilis complete genom					
[NT:similar to acetyl-CoA C-acetyltr					
>gp:[GI:e324998:g2226255] [LN:BSY140					
[OR:Bacillus subtilis] [DB:genpept-bdegrees: aprE to comK.] [NT:similari					
[LE:3788] [RE:4882] [DI:complement]	.ty to a	cerar-	COAuacei	cyrtransfera	ises (Thil,)
[LEDVO / OF [ARE TOOL] [DITOMPTOMOTE]				 	
ORF Name	NTID	AAID	NT	AA score	probability
A17503000987 39635 £2 316	1618	5390	LengthI 1245		7.4e-98
Description			التتال		
sp:[LN:YKGC ECOLI] [AC:P77212] [GN:Y	מאמכו (סו	ם. בכרטו	בסדרטדא	COLLI [DE.I	NTEDCENIC DECION
[SP:P77212] [DB:swissprot] >pir:[LN:				_	_
reductase,:ykgC protein] [GN:ykgC]				_	-
>gp:[GI:g1657503] [LN:ECU73857] [AC:					
[DE:Escherichia coli chromosome minu					
reductase] [LE:26473] [RE:27825] [DI	:compler	ment] :	gp:[GI:	:g1786495] [LN:AE000137]
[AC:AE000137:U00096] [PN:putative ox					
classified] [OR:Escherichia coli] [D					
section 27 of 400 of the completegen	ome.] [1	NT:£450	o; 35 po	ct identical	(29 gaps) to 430

residues of] [LE:6292] [RE:7644] [DI:complement]

			NT AA	
ORF Name	NTID	AAID	LengthLength score	probability
A17503000987_4031952_c2_775	1619	5391	501 166 545 1	3e-52
Description	,			
pir:[LN:D69695] [AC:D69695] [PN:rik [CL:Escherichia coli ribosomal prote >gp:[GI:d1009473:g786163] [LN:BACRPI [OR:Bacillus subtilis] [SR:Bacillus [DE:Bacillus subtilis genes for ribocds.] [LE:347] [RE:847] [DI:direct] [AC:Z99104:AL009126] [PN:ribosomal r [DB:genpept-bct1] [DE:Bacillus subtito213080.] [SP:P42923] [LE:120057]	ein L10] L2] [AC: subtili csomal p >gp:[GI crotein ilis com	[OR:Ba D50303] s (stra roteins :e11820 L10 (BI	acillus subtilis] [I [PN:Ribosomal Protain:168) DNA] [DB:ge L1, L10 and L12,pa D37:g2632371] [LN:BS L5)] [GN:rplJ] [OR:E	DB:pir2] tein L10] [GN:rplJ] enpept-bct1] artial and complete SUB0001] Bacillus subtilis]
ORF Name	NTID	AAID	NT AA LengthLength	probability
AI7503000987_40676_c1_644	1620	5392	183 60	
Description				
NO-HIT				The second secon
ORF Name	NTID	AAID	NT AA LengthLength	probability
A17503000987_4078305_c3_927	1621	5393	126 41	
Description				
NO-HIT				
ORF Name	NTID	AAID	NT AA LengthLength score	probability
A17503000987_4078375_f1_6	1622	5394	129 42	
Description				
NO-HIT				
ORF Name	NTID	AAID	NT AA LengthLength score	probability
AI7503000987_4078518_f1_143	1623		843 280 521 4	.6e-50
Description			_	_
sp:[LN:YXEK_BACSU] [AC:P54950] [GN:Y49.3 KD PROTEIN IN IDH-DEOR INTERGEN				
>pir: [LN:E70075] [AC:E70075] [PN:mo				
[CL:nitrilotriacetate monooxygenase]			-	
>gp:[GI:e1184677:g2636498] [LN:BSUB0				
[OR:Bacillus subtilis] [DB:genpept-k				
21 of 21): from 3999281to 4214814.]				
[LE:60328] [RE:61653] [DI:complement [AC:D45912] [GN:yxeK] [OR:Bacillus s				
(Marburg 168; trpC2)) DNA] [DB:genpe				
between the iol and hut operon, parti				
product of Rhodococcus sp.] [LE:9823				_

 $\underline{A}\underline{A}$ NT ORF Name AAID NTID score probability LengthLength AI7503000987 4181527 cl 720 1624 5396 447 148 Description sp:[LN:V17 BPT7] [AC:P03781] [GN:1.7] [OR:BACTERIOPHAGE T7] [DE:GENE 1.7 PROTEIN] [SP:P03781] [DB:swissprot] >pir:[LN:W1BP77] [AC:G43002:G43004:S42296:A04406] [PN:qene 1.7 protein] [GN:1.7] [CL:phage T7 gene 1.7 protein] [OR:phage T7] [DB:pir1] [MP:20.42-21.89] >gp:[GI:g15512] [LN:PET7XX] [AC:V01127] [OR:Bacteriophage T7] [DB:genpept-phg] [DE:Left end of bacteriophage T7 genome. Includes the reading frames of the genes 0.3, 0.4, 0.5, 0.6, 0.65, 0.7, 1, 1.1, 1.2, 1.3 (early proteins) and 1.4, 1.5, 1.6, 1.7, 2, 2.5, 2.8, 3, 3.5, 4A and 4B(late proteins). Gene 1 is the T7 RNA polymerase.] [NT:1.7 protein] [SP:P03781] [LE:8166] [RE:8756] [DI:direct] >gp:[GI:g15576] [LN:T7CG] [AC:V01146:J02518:X00411] [OR:Bacteriophage T7] [DB:genpept-phg] [DE:Genome of bacteriophage T7.] [NT:gene 1.7] [SP:P03781] [LE:8166] [RE:8756] [DI:direct] AΑ ORF Name NTID AAID probability score LengthLength AI7503000987 4181577 c3 955 1251 3.9e-441625 416 465 Description pir: [LN:H70427] [AC:H70427] [PN:replicative DNA helicase] [GN:dnaB] [CL:phage P22 gene 12 protein] [OR:Aquifex aeolicus] [DB:pir2] >qp:[GI:q2983861] [LN:AE000742] [AC:AE000742:AE000657] [PN:replicative DNA helicase] [GN:dnaB] [OR:Aquifex aeolicus] [DB:genpept-bct2] [DE:Aquifex aeolicus section 74 of 109 of the complete genome.] [LE:2249] [RE:3655] [DI:direct] NT AA ORF Name NTID AAID score probability LengthLength A17503000987 4329063 c3 900 1626 1245 1786 4.1e-184 Description gp:[GI:d1039005:g4512402] [LN:AB017508] [AC:AB017508] [GN:tufA] [OR:Bacillus halodurans [SR:Bacillus halodurans (strain:C-125) DNA] [DB:genpept-bct1] [DE:Bacillus halodurans C-125 genomic DNA, 32 kb fragment, completecds.] [NT:tufA homologue (identity of 91% to B. subtilis)] [LE:11150] [RE:12340] [DI:direct] NT AΑ ORF Name NTID AAID score probability LengthLength A17503000987 4334510 c2 822 1627 5399 135 Description NO-HIT NT AΑ ORF Name NTID AAID score probability LengthLength A17503000987_4398453_f3_561 5400 158 1628 477 4.5e-24 279 Description sp:[LN:NTAA CHEHE] [AC:P54989] [GN:NTAA:NMOA] [OR:CHELATOBACTER HEINTZII] [EC:1.14.13.-] [DE:MONOOXYGENASE COMPONENT A) (NTA-MO A)] [SP:P54989] [DB:swissprot] >gp:[GI:g1119211] [LN:CBYNMOB] [AC:L49438] [GN:nmoA] [FN:component A of NTA monooxygenase] [OR:Chelatobacter heintzii] [DB:genpept-bct1] [DE:Chelatobacter heintzii NTA monooxygenase component B (nmoB), NTAmonooxygenase component A (nmoA), regulatory protein (nmoR) andtransposase (nmoT) genes, complete cds.] [LE:1323] [RE:2684] [DI:direct] >gp:[GI:g1480205] [LN:CHU39411] [AC:U39411] [PN:NTA monooxygenase component A] [GN:ntaA] [OR:Chelatobacter heintzii] [SR:Chelatobacter heintzii strain=ATCC 29600] [DB:genpept-bct1] [DE:Chelatobacter heintzii nitrilotriacetate monooxygenase

genes, putative regulatory protein (ntaR), NTA monooxygenase component A(ntaA) and component B (ntaB) genes, complete cds.] [LE:1123] [RE:2484] [DI:complement]

ORF Name	NTID AAID NT AA score probability
A17503000987_446062_c1_748	
Description	
SEQUENCE-LIKE ELEMENT IS431MEC] [SP: [AC:S12093:JU0116] [PN:probable traps] >gp: [GI:g46602] [LN:SAIS431M] [AC:X5] [DB:genpept-bct1] [DE:S. aureus IS43] [NT:putative transposase (AA 1 - 224 >gp: [GI:e1237900:g2791991] [LN:SAMEC [OR:Staphylococcus aureus] [DB:genpemecI genes and ORF168, ORF142,ORF44, [RE:8770] [DI:direct] >gp: [GI:d10460] [PN:transposase for insertion sequent [SR:Staphylococcus aureus (strain:N3] [DE:Staphylococcus aureus genes, med [LE:48054] [RE:48728] [DI:direct] >g [PN:transposase for insertion sequent [SR:Staphylococcus aureus (strain:N3)]	ETAPHYLOCOCCUS AUREUS] [DE:TRANSPOSASE FOR INSERTION Pl9380] [DB:swissprot] >pir:[LN:S12093] ansposase] [OR:Staphylococcus aureus] [DB:pir2] ansposase] [OR:Staphylococcus aureus] almec gene associated with methicillin resistance.] [DB:P:P19380] [LE:272] [RE:946] [DI:direct] and [DE:Staphylococcus aureus mecA, mecR1, and [DE:Staphylococcus aureus mecA, mecR1, and ORF145 and ORF224.] [NT:ORF224] [LE:8096] align="1">[DE:Staphylococcus aureus mecA, mecR1, and [DB:Staphylococcus aureus] are-like element] [OR:Staphylococcus aureus] are-like element] [OR:Staphylococcus aureus] aregion, partial and complete cds.] [NT:ORF N062] are-like element] [OR:Staphylococcus aureus] are-like element] [OR:St
ORF Name A17503000987_4725006_c3_981	NTID AAID NT AA score probability 1630 5402 537 178
Description NO-HIT	
ORF Name	NTID AAID NT AA score probability
A17503000987_4727203_c3_925	1631 5403 369 122 86 0.032
structural protein 1] [OR:Mus muscul spermatocyte cDNA t] [DB:genpept-rod	[AC:D14849] [PN:meiosis-specific nuclear us] [SR:Mus mucsulus (strain ddY) testis pachytene [DE:Mouse mRNA for meiosis-specific nuclear cds.] [LE:180] [RE:1655] [DI:direct]
ORF Name	NTID AAID LengthLength score probability
AI7503000987_4735833_t2_359	1632 5404 711 236 464 5.0e-44
Description	
KD PROTEIN IN SERS-DNAZ INTERGENIC R [AC:S66045:G69736] [PN:deoxypurine [CL:Lactobacillus acidophilus deoxya >gp:[GI:d1005793:g467405] [LN:BAC180 [SR:Bacillus subtilis (sub_species:M subtilis DNA, 180 kilobase region of [DI:complement] >gp:[GI:e1181948:g26 [FN:unknown] [OR:Bacillus subtilis]	[AAG] [OR:BACILLUS SUBTILIS] [DE:HYPOTHETICAL 24.1 EGION] [SP:P37530] [DB:swissprot] >pir:[LN:S66045] kinase subunit homolog yaaG] [GN:yaaG] denosine kinase] [OR:Bacillus subtilis] [DB:pir2] [AC:D26185] [PN:unknown] [OR:Bacillus subtilis] [DE:Burg, strain:168) DNA] [DB:genpept-bct1] [DE:B. replication origin.] [LE:86746] [RE:87369] 32282] [LN:BSUB0001] [AC:Z99104:AL009126] [GN:yaaG] [DB:genpept-bct1] [DE:Bacillus subtilis complete 13080.] [NT:similar to deoxypurine kinase subunit] I:complement]

ORF Name	NTID AAID NT AA score probability
AI7503000987_4741068_c3_967	1633 5405 609 202 210 4.1e-17
Description	
gp:[GI:e247154:g1926352] [LN:LBPHIG1	E] [AC:X98106] [PN:minor capsid protein] e] [DB:genpept-phg] [DE:Lactobacillus bacteriophage 554] [RE:25168] [DI:complement]
ORF Name	NTID AAID NT AA score probability
AI7503000987_4741077_c3_952	1634 5406 687 228 238 4.5e-20
Description	
	E] [AC:X98106] [GN:Rorf242] [OR:Bacteriophage cillus bacteriophage phig1e complete genomic DNA.]
ORF Name	NTID AAID NT AA score probability
AI7503000987_4876967_c3_913	1635 5407 579 192 168 1.2e-12
Description	
[AC:Z99108:AL009126] [GN:yhdA] [FN:u [DE:Bacillus subtilis complete genom [NT:similar to hypothetical proteins >gp:[GI:e1182935:g2633269] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-b6 of 21): from 999501 to1209940.] [N [RE:10926] [DI:direct] >gp:[GI:e3249 [PN:hypothetical protein] [GN:yhdA] [DE:Bacillus subtilis chromosomal DN	:[GI:e1182923:g2633257] [LN:BSUB0005] nknown] [OR:Bacillus subtilis] [DB:genpept-bct1] e (section 5 of 21): from 802821 to1011250.]] [LE:207082] [RE:207606] [DI:direct] 006] [AC:Z99109:AL009126] [GN:yhdA] [FN:unknown] ct1] [DE:Bacillus subtilis complete genome (section T:similar to hypothetical proteins] [LE:10402] 46:g2226142] [LN:BSY14079] [AC:Y14079] [OR:Bacillus subtilis] [DB:genpept-bct1] A, region 75 degrees: glpPFKDoperon and ypothetical protein YIEF from] [LE:10050]
ORF Name	NTID AAID NT AA probability
A17503000987_4882760_c3_1001	1636 5408 183 60
Description NO-HIT	
ORF Name	NTID AAID NT AA score probability
A17503000987_4886075_£3_559	1637 5409 678 225 794 5.4e-79
Description	
KD PROTEIN IN SERS-DNAZ INTERGENIC R [AC:S66044:F69736] [PN:deoxypurine [CL:Lactobacillus acidophilus deoxya >gp:[GI:d1005792:g467404] [LN:BAC180 [SR:Bacillus subtilis (sub_species:M subtilis DNA, 180 kilobase region of [DI:complement] >gp:[GI:e1181947:g26 [FN:unknown] [OR:Bacillus subtilis]	AAF] [OR:BACILLUS SUBTILIS] [DE:HYPOTHETICAL 25.4 EGION] [SP:P37529] [DB:swissprot] >pir:[LN:S66044] kinase subunit homolog yaaF] [GN:yaaF] denosine kinase] [OR:Bacillus subtilis] [DB:pir2] K] [AC:D26185] [PN:unknown] [OR:Bacillus subtilis] arburg, strain:168) DNA] [DB:genpept-bct1] [DE:B. replication origin.] [LE:86096] [RE:86749] 32281] [LN:BSUB0001] [AC:Z99104:AL009126] [GN:yaaF] [DB:genpept-bct1] [DE:Bacillus subtilis complete 13080.] [NT:similar to deoxypurine kinase subunit] I:complement]

NT AΑ ORF Name AAID NTID probability LengthLength A17503000987 4892127 cl 746 1638 5410 2706 901 7.6e-160 Description pir: [LN:H69780] [AC:H69780] [PN:antibiotic transport-associated protein homolog ydfJ] [GN:ydfJ] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:d1020130:g1881350] [LN:AB001488] [AC:AB001488] [GN:ydfJ] [OR:Bacillus subtilis] [SR:Bacillus subtilis (strain:168) DNA] [DB:genpept-bct1] [DE:Bacillus subtilis genome sequence, 148 kb sequence of the regionbetween 35 and 47 degree.] [NT:PROBABLE TRANSPORT PROTEIN, SIMILAR TO ANTIBIOTIC] [LE:122935] [RE:125109] [DI:direct] >qp:[GI:e1182509:q2632843] [LN:BSUB0003] [AC:Z99106:AL009126] [GN:ydfJ] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 3 of 21): from 402751 to611850.] [NT:similar to antibiotic transport-associated protein] [LE:186681] [RE:188855] [DI:direct] ORF Name AAID NTID probability score LengthLength AI7503000987_4901703 c2 792 1639 5411 510 169 0.0031 Description pir: [LN:D69887] [AC:D69887] [PN:conserved hypothetical protein ynaD] [GN:ynaD] [OR:Bacillus subtilis] [DB:pir2] >qp: [GI:q1750115] [LN:BSU66480] [AC:U66480] [PN:YnaD] [GN:ynaD] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis SpoVK (spoVK), YnbA (ynbA), YnbB (ynbB), GlnR(glnR), glutamine synthetase (glnA), YnaA (ynaA), YnaB (ynaB), YnaC(ynaC), YnaD (ynaD), YnaE (ynaE), YnaF (ynaF), YnaG (ynaG), YnaH(ynaH), YnaI (ynaI), YnaJ (ynaJ), xylan beta-1,4-xylosidase (xynB),xylose repressor (xylR), xylose isomerase (xylA), xylulose kinase(xylB), YncB (yncB), YncC (yncC), YncD (yncD) and YncE (yncE)genes, complete cds.] [LE:9169] [RE:9681] [DI:direct] >gp:[GI:e1183411:g2634136] [LN:BSUB0010] [AC:Z99113:AL009126] [GN:ynaD] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 10 of 21): from 1781201to 2014980.] [NT:similar to hypothetical proteins] [LE:101220] [RE:101732] [DI:direct] NT AΑ NTID AAID score probability LengthLength 1640 5412 1068 355 183 1.0e-11 Description gp:[GI:g3264834] [LN:AF072541] [AC:AF072541] [PN:xylitol dehydrogenase] [GN:xdh] [FN:xylose utilisation] [OR:Galactocandida mastotermitis] [DB:qenpept-pln2]

ORF Name A17503000987_4946962 c3 907

[EC:1.1.1.9] [DE:Galactocandida mastotermitis xylitol dehydrogenase (xdh) gene, complete cds.] [NT:a member of the medium chain dehydrogenase] [LE:301:373] [RE:312:1422] [DI:directJoin]

NT AΑ ORF Name NTID AAID score probability Length Length A17503000987 4964677 c2 832 1641 4445413 147 4.3e-22

Description

pir:[LN:T00179] [AC:T00179] [PN:ssDNA-binding protein] [OR:Staphylococcus aureus phage phi PVL] [DB:pir3] >gp:[GI:d1032880:g3341953] [LN:AB009866] [AC:AB009866] [PN:ssDNA binding protein] [OR:bacteriophage phi PVL] [SR:bacteriophage phi PVL (specific host:Staphylococcus aureus ATC] [DB:genpept-phg] [DE:Bacteriophage phi PVL proviral DNA, complete sequence.] [NT:orf 45] [LE:34657] [RE:35127] [DI:direct]

ORF Name NTID probability A17503000987_5080092_c2_809 5414 744 1642 6.5e-51 Description sp:[LN:YBFT BACSU] [AC:031458] [GN:YBFT] [OR:BACILLUS SUBTILIS] [DE:HYPOTHETICAL 27.3 KD PROTEIN IN GLTP-CWLJ INTERGENIC REGION] [SP:031458] [DB:swissprot] >pir:[LN:E69750] [AC:E69750] [PN:glucosamine-6-phosphate isomerase homolog ybfT] [GN:ybfT] [CL:glucosamine-6-phosphate isomerase] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:d1034099:g3599658] [LN:AB006424] [AC:AB006424] [GN:ybfT] [OR:Bacillus subtilis] [SR:Bacillus subtilis (strain:168) DNA] [DB:genpept-bct1] [DE:Bacillus subtilis genomic DNA, 70 kb region between 17 and 23degree.] [LE:59813] [RE:60562] [DI:complement] >qp:[GI:e1182188:g2632522] [LN:BSUB0002] [AC:Z99105:AL009126] [GN:ybfT] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 2 of 21): from 194651 to415810.] [NT:similar to glucosamine-6-phosphate isomerase] [SP:031458] [LE:62162] [RE:62911] [DI:complement] NT AΑ ORF Name NTID AAID probability score LengthLength AI7503000987_5125076_f3_531 810 2.3e-87 1643 Description sp:[LN:YWFI BACSU] [AC:P39645] [GN:YWFI:IPA-87R] [OR:BACILLUS SUBTILIS] [DE:HYPOTHETICAL 29.5 KD PROTEIN IN ROCC-PTA INTERGENIC REGION] [SP:P39645] [DB:swissprot] >pir:[LN:S39742] [AC:S39742:B70056] [PN:ywfI protein:hypothetical protein ipa-87r] [GN:ywfI] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:g414011] [LN:BSGENR] [AC:X73124] [GN:ipa-87r] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:B.subtilis genomic region (325 to 333).] [SP:P39645] [LE:90200] [RE:90964] [DI:complement] >qp:[GI:e1186267:q2636303] [LN:BSUB0020] [AC:Z99123:AL009126] [GN:ywfI] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 20 of 21): from 3798401to 4010550.] [NT:alternate gene name: ipa-87r; similar to] [SP:P39645] [LE:66995] [RE:67759] [DI:direct] AA ORF Name <u>sco</u>re NTID AAID probability LengthLength AI7503000987_5251588_c1_747 1644 5416 582 193 0.0010 Description gp:[GI:g1079814] [LN:S79230] [AC:S79230] [PN:BM1P1] [GN:CYP106] [OR:Bacillus megaterium] [DB:genpept-bct2] [DE:CYP106=BM1P2 orf...CYP106=P450BM-1 orf {regulatory regions [Bacillus megaterium, mRNA Partial, 3 genes, 1400 nt].] [NT:positive transcription factor involved in] [LE:763] [RE:1059] [DI:complement] NT AA ORF Name NTID AAID probability LengthLength AI7503000987_5258515_c3_965 1645 1314 2.1e-132 5417 437 Description

sp:[LN:TERL BPSPP] [AC:P54308] [GN:2] [OR:BACTERIOPHAGE SPP1] [DE:TERMINASE LARGE SUBUNIT (G2P)] [SP:P54308] [DB:swissprot] >pir:[LN:S24451] [AC:S24451] [PN:terminase] [OR:phage SPP1] [DB:pir2] >gp:[GI:e244468:g2764840] [LN:BSPP1GENM] [AC:X97918] [OR:Bacteriophage SPP1] [DB:genpept-phg] [DE:Bacteriophage SPP1 complete nucleotide sequence.] [NT:gene 2] [SP:P54308] [LE:307] [RE:1575] [DI:direct] >gp:[GI:q15466] [LN:NCSPP116] [AC:X56064:S35313] [PN:terminase] [GN:SPP1 gene 2] [OR:Bacteriophage SPP1] [DB:genpept-phg] [DE:B. subtilis bacteriophage SPP1 genes for terminase and portalprotein.] [SP:P54308] [LE:633] [RE:1901] [DI:direct]

ORF Name	NTID	AAID	NT AA score probability
A17503000987_5370450_±3_432	1646	5418	1284 427 305 3.6e-27
Description			
pir: [LN:A69256] [AC:A69256] [PN:hypfulgidus] [DB:pir2] >gp:[GI:g2650605fulgidus predicted coding region AF0 [DB:genpept-bct2] [DE:Archaeoglobus	[LN:AE 049] [GN fulgidus	00110 N:AF00 s sect	3] [AC:AE001103:AE000782] [PN:A.
ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000987_56626_c2_855	1647	5419	336 111 84 0.0020
	enpept-b oprotein	oct1] n (LP)	
ORF Name	NTID	AAID	NT AA score probability
AI7503000987_587811_£1_125	1648	5420	345 114
Description NO-HIT			
ORF Name	NTID	AAID	NT AA score probability
A17503000987_6056625_c1_755	1649	5421	258 85
Description NO-HIT			
ORF Name	NTID	AAID	NT AA LengthLength score probability
AI7503000987_6149152_c3_884	1650	5422	1389 462 1543 2.3e-158
] [CL:DNA repair protein sms] [OR:Ba >gp:[GI:d1005863:g467475] [LN:BAC180 [SR:Bacillus subtilis (sub_species:M subtilis DNA, 180 kilobase region of [DI:direct] >gp:[GI:e1182020:g263235 repair protein homolog] [GN:sms] [OR	TEIN SMS 69709] cillus s K] [AC:D arburg, replica 4] [LN:B :Bacillu of 21):	PN:DI ubtil: 26185; strain tion of SUB000 s subf	LOG)] [SP:P37572] [DB:swissprot] NA repair protein homolog sms] [GN:sms is] [DB:pir2]] [PN:unknown] [OR:Bacillus subtilis] n:168) DNA] [DB:genpept-bct1] [DE:B. origin.] [LE:169695] [RE:171071] O1] [AC:Z99104:AL009126] [PN:DNA tilis] [DB:genpept-bct1] [DE:Bacillus 1 to213080.] [NT:alternate gene name:
ORF Name A17503000987_6250050_f2_233 Description		<u>AAID</u> 5423	NT AA LengthLength score probability 147 48

NO-HIT

NT AA ORF Name AAID score probability NTID LengthLength A17503000987 6406337 ±3 448 1652 210 69 Description NO-HIT NTAA ORF Name NTID AAID probability score LengthLength A17503000987 6407136_c2_840 1653 5425 1455 Description sp:[LN:SIZ BPSPP] [AC:P54309] [GN:6:SIZ] [OR:BACTERIOPHAGE SPP1] [DE:PORTAL PROTEIN (PORTAL VERTEX PROTEIN) (GP6)] [SP:P54309] [DB:swissprot] >pir:[LN:S21805] [AC:S21805:S24455:S36725] [PN:portal protein:gene 6 protein] [GN:6] [OR:phage SPP1] [DB:pir2] >gp:[GI:e244702:g2764847] [LN:BSPP1GENM] [AC:X97918] [OR:Bacteriophage SPP1] [DB:genpept-phg] [DE:Bacteriophage SPP1 complete nucleotide sequence.] [NT:gene 6] [SP:P54309] [LE:2334] [RE:3845] [DI:direct] >gp:[GI:g15470] [LN:NCSPP116] [AC:X56064:S35313] [PN:portal protein] [GN:SPP1 gene 6] [OR:Bacteriophage SPP1] [DB:qenpept-phq] [DE:B. subtilis bacteriophage SPP1 genes for terminase and portalprotein.] [SP:P54309] [LE:2660] [RE:4171] [DI:direct] ORF Name NTID score probability Length Length AI7503000987_6443763_c2_769 408 2.0e-26 1654 5426 135 298 Description pir:[LN:C69742] [AC:C69742] [PN:conserved hypothetical protein yazC] [GN:yazC] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1182028:g2632362] [LN:BSUB0001] [AC:Z99104:AL009126] [GN:yazC] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 1 of 21): from 1 to213080.] [NT:similar to hypothetical proteins] [LE:114851] [RE:115282] [DI:direct] NT ORF Name NTID score AAID probability LengthLength A17503000987 661062 c1 641 1655 5427 1440 1568 5.2e-161 Description sp:[LN:SYC BACSU] [AC:Q06752] [GN:CYSS:SPNA] [OR:BACILLUS SUBTILIS] [EC:6.1.1.16] [DE: (CYSRS)] [SP:Q06752] [DB:swissprot] >pir: [LN:C53402] [AC:C53402:S44447:S66123:B69612] [PN:cysteine--tRNA ligase, cysS:cysteinyl-tRNA synthetase cysS] [GN:cysS] [CL:cysteine--tRNA ligase] [OR:Bacillus subtilis] [EC:6.1.1.16] [DB:pir1] >gp:[GI:d1005870:g467482] [LN:BAC180K] [AC:D26185] [PN:cysteinyl-tRNA synthetase] [GN:cysS] [OR:Bacillus subtilis] [SR:Bacillus subtilis (sub species: Marburg, strain: 168) DNA] [DB:genpept-bct1] [DE:B. subtilis DNA, 180 kilobase region of replication origin.] [LE:177049] [RE:178449] [DI:direct] >gp:[GI:g289284] [LN:BACGLUSYN] [AC:L14580] [PN:cysteinyl-tRNA synthetase] [OR:Bacillus subtilis] [SR:Bacillus subtilis (strain 168T) DNA] [DB:genpept-bct1] [DE:Bacillus subtilis glutamyl-tRNA transferase (gltX), serineacetyltransferase (cysE), and cysteinyl-tRNA synthetase (cysS)genes, complete cds's.] [LE:4109] [RE:5509] [DI:direct] >gp:[GI:g499303] [LN:BSCTS] [AC:X73989] [PN:cysteine--tRNA ligase] [GN:SPNA/CYSS] [OR:Bacillus subtilis] [DB:genpept-bct1] [EC:6.1.1.16] [DE:B.subtilis gene for cysteinyl-tRNA synthetase.] [SP:Q06752] [LE:321] [RE:1721] [DI:direct] >gp:[GI:e1182027:g2632361] [LN:BSUB0001] [AC:Z99104:AL009126] [PN:cysteinyl-tRNA synthetase] [GN:cysS] [OR:Bacillus subtilis] [DB:genpept-bct1] [EC:6.1.1.16]

[DE:Bacillus subtilis complete genome (section 1 of 21): from 1 to213080.]
[NT:alternate gene name: snpA] [SP:Q06752] [LE:113447] [RE:114847] [DI:direct]

ORF Name	NTID	AAID	NT AA score probability
AI7503000987 6725707_c2_770			Lengthbength
Description	1656	5428	552 183 327 1.7e-29
sp:[LN:YACP_BACSU] [AC:P37574] [GN:YACP_BACSU] [AC:P37574] [GN:YACP_BACSU] [AC:P37574] [GN:YACP_BACSU] [PN:conserved hysubtilis] [DB:pir2] >gp:[GI:d1005872] [OR:Bacillus subtilis] [SR:Bacillus [DB:genpept-bct1] [DE:B. subtilis DN [LE:179624] [RE:180136] [DI:direct] [AC:Z99104:AL009126] [GN:yacP] [FN:VACP] [FN:VACP] [FN:VACP]	7574] [D. Potheti 2:g46748 subtili NA, 180 >gp:[GI unknown] ne (sect	B:swiss cal pro 4] [LN s (sub_ kilobas :e11820 [OR:Ba ion 1 c	rotein yacP] [GN:yacP] [OR:Bacillus I:BAC180K] [AC:D26185] [PN:unknown] p_species:Marburg, strain:168) DNA] ase region of replication origin.] compared to the subtilis of 21): from 1 to 213080.] [NT:similar
ORF Name	NTID	AAID	NT AA score probability
AI7503000987_6757338_c3_906	1657	5429	1089 362 1129 1.7e-114
Description	·		
	3:g35996 subtilia 70 kb reg 2191:g26 subtilia From 1940	62] [L1 s (stra gion be 32525] s] [DB: 651 to4	N:AB006424] [AC:AB006424] [GN:ybgE] rain:168) DNA] [DB:genpept-bct1] retween 17 and 23degree.] [LE:62005]
ORF Name	NTID	AAID	NT AA score probability
A17503000987_6932750_f2_348	1658	5430	
Description pir:[LN:C69906] [AC:C69906] [PN:hypsubtilis] [DB:pir2] >gp:[GI:e1185419] [GN:yojF] [FN:unknown] [OR:Bacillus complete genome (section 11 of 21):[DI:complement] >gp:[GI:g3169322] [I [FN:unknown] [OR:Bacillus subtilis] (yojA), YojB (yojB), YojC (yojC), YoYojH (yojH), YojI(yojI), YojJ (yojJ) YojN(yojN), and YojO (yojO) genes, Calle:3518] [RE:3868] [DI:direct]	subtilis from 200 N:AF0261 [DB:genp jD(yojD) , YojK	40] [LNs] [DB: 00171tc 147] [Apept-bo), YojE (yojK),	N:BSUB0011] [AC:Z99114:AL009126] e:genpept-bct1] [DE:Bacillus subtilis o 2207900.] [LE:121397] [RE:121747] AC:AF026147] [PN:YojF] [GN:yojF] ct2] [DE:Bacillus subtilis YojA E (yojE), YojF (yojF), YojG (yojG), , YojL (yojL), YojM (yojM),
ORF Name	NTID	AAID	NT AA LengthLength score probability

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
AI7503000987_7068751_c3_984	1660	5432	945	314		4.6e-98
Description		L	لــــال		L	
sp:[LN:IOLS_BACSU] [AC:P46336] [GN:I (VEGETATIVE PROTEIN 147) (VEG147)] [AC:D69646] [PN:myo-inositol catabo protein YPL088w] [OR:Bacillus subtil [LN:AB005554] [AC:AB005554:D45242:D3 subtilis (strain:BGSC 1A1) DNA] [DB:kb region between gnt and ioloperons [LE:32539] [RE:33471] [DI:complement [AC:Z99124:AL009126] [GN:iolS] [OR:B subtilis complete genome (section 21 gene name: yxbF; myo-inositol catabo	SP:P463 Plism io is] [DB 1629] [genpept s.] [NT:] >gp:[acillus of 21)	36] [D: ls] [G: :pir2] GN:iol: -bct1] plausil GI:e11: subti: : from	B:swiss N:iolS >gp:[G S] [OR: [DE:Ba bly inv 84703:g lis] [D 399928	sprot]] [CL: GI:d102:Bacill acillus volved g263652 DB:genp	>pir:[conser 2457:g us subtiin ino 4] [LN ept-bc 14814.	LN:D69646] rved hypothetical 904205] tilis] [SR:Bacillus lis genomic DNA, 36 sitol catabolism] T:BSUB0021] tt] [DE:Bacillus] [NT:alternate
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
AI7503000987_7160287_c2_805	1661	5433	126	41		
Description			لــــــــــا <i>ل</i> ـ	L-,	l	
NO-HIT						
ORF Name AI7503000987_7164191_c2_798	NTID 1662	<u>AAID</u> 5434	NT Length	AA Length 49	score	probability
Description			J	L		
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000987_7229680_c1_634	1663	5435	492	163	377	8.3e-35
Description						
sp:[LN:CTSR_BACSU] [AC:P37568] [GN:CREGULATOR CTSR] [SP:P37568] [DB:swis [PN:transcription repressor of class subtilis] [DB:pir2] >gp:[GI:d1005859] [OR:Bacillus subtilis] [SR:Bacillus [DB:genpept-bct1] [DE:B. subtilis DN [LE:165048] [RE:165512] [DI:direct] [AC:Z99104:AL009126] [PN:transcripti of class III stress genes] [OR:Bacil subtilis complete genome (section 1 yacG] [SP:P37568] [LE:101446] [RE:10	sprot] :g46747 :subtilia A, 180 >gp:[GI onal red lus subtof 21):	>pir:[] ress ge 1] [LN s (sub_ kilobas :e11820 gulaton tilis] from 1	LN:S661 enes ct :BAC180 _specie se regi D16:g26 r] [GN: [DB:ge L to213	12] [A sR] [G K] [AC s:Marb on of [32350] ctsR]	C:S661 N:ctsR :D2618 urg, s replica [LN:Barreneral]	12:D69610]] [OR:Bacillus 5] [PN:unknown] train:168) DNA] ation origin.] SUB0001] gative regulation [DE:Bacillus
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
AI7503000987_7275263_c1_733	1664	5436	441	146	158	1.3e-11
Description gp:[GI:e244713:g2764865] [LN:BSPP1GE [DB:genpept-phg] [DE:Bacteriophage S [LE:10063] [RE:10467] [DI:direct]						

ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	score	probability
A17503000987_782590_£1_114	1665	5437	171	56	56	0.044
Description gp:[GI:e289995:g1764015] [LN:CICOS41] [DB:genpept-inv1] [DE:Ciona intesting similarity to membrane transport proff [RE:13190:13703:14681] [DI:complement)	alis DN teins]	A sequ	ence fr	om cos	smid C	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000987_783425_c1_675	1666	5438		54]	
Description NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> <u>Length</u>	<u>AA</u> Length	score	probability
A17503000987_818942_c3_989	1667	5439	477	158	173	3.4e-17
BETAINE-BINDING PROTEIN PRECURSOR] [8 [AC:I40537:F69669] [PN:glycine beta:protein) opuAC precursor] [GN:opuAC] protein) opuAC precursor] [GN:opuAC] protein) opuAC precursor] [GN:opuAC] [FN:BSUB06] [GN:opuAC] [FN:OR:Bacillus subtilis] [DB:genpept-bc2 of 21): from 194651 to415810.] [SP:pg:[GI:d1009569:g1805372] [LN:D50455] precursor] [GN:opuAC] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis] [DB:genpept-bct1] [DB:Bacillus subtilis] [DB:genpept-bct1] [OR:Bacillus subtilis] [DB:genpept-bct1] [DB:gen	ine ABC [OR:Bactorial Color of the color of	trans; acillu: C:Z991; e beta: E:Baci:] [LE:: D50453;] [SR:: s DNA: s DNA: [RE ne-bing taine-l	porter s subti 05:AL00 ine tra llus su 128023] [PN:g Bacillu for 25- :3207] ding pr llus su binding	(glyci lis] [9126] Insport btilis [RE:1 lycine Is subt 36 deg [DI:di Totein btilis	Ine before the precure of ATPass	taine-binding r2] lycine betaine ABC oprotection)] lete genome (section] [DI:direct] in-binding protein (strain:168 trpC2) egion containing >gp:[GI:g984805] rsor] [GN:opuAC] se (opuAA),
ORF Name	NTID	AAID	<u>NT</u> Length:	<u>AA</u> Length	score	probability
AI7503000987_819575_c2_850	1668	5440		618		1.4e-08
Description gp:[GI:e247163:g1926361] [LN:LBPHIGING phig1e] [DB:genpept-phg] [DE:Lactobace [LE:13476] [RE:14594] [DI:complement]	cillus 1					
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000987_821012_c1_664	1669	5441	1725	574	469	6.8e-44
Description pir: [LN:S49240] [AC:S49240] [PN:hypo [OR:Haemophilus influenzae] [DB:pir2] [OR:Haemophilus influenzae] [DB:genpe capsulation locus region II DNA.] [NT	>gp:[0	GI:g54'	7513] [:Haemop	LN:HIA hilus	CAPIII influe	D] [AC:Z37516] enzae serotype a
ORF Name	NTID	AAID	<u>NT</u> Lengthl	<u>AA</u> Length	score	probability
A17503000987_833125_±3_493	1670	5442		94		
Description NO-HIT					•	

ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000987 892842_c3_926	1671	5443	
Description			
sp:[LN:YWDK_BACSU] [AC:P39619] [GN:Y [DE:HYPOTHETICAL 12.0 KD PROTEIN IN [DB:swissprot] >pir:[LN:S39716] [AC: protein ipa-61d] [GN:ywdK] [OR:Baci [LN:BSGENR] [AC:X73124] [GN:ipa-61d] [DE:B.subtilis genomic region (325 t [DI:direct] >gp:[GI:e1186292:g263632 [FN:unknown] [OR:Bacillus subtilis]	UNG-ROCA S39716:I llus suk [OR:Bac to 333).] 8] [LN:I [DB:genry	A INTE F70054 otilis cillus [SP: BSUB00 pept-b	RGENIC REGION] [SP:P39619]] [PN:ywdK protein:hypothetical] [DB:pir2] >gp:[GI:g413985] subtilis] [DB:genpept-bct1] P39619] [LE:63169] [RE:63510] 20] [AC:Z99123:AL009126] [GN:ywdK] ct1] [DE:Bacillus subtilis complete .] [NT:alternate gene name: ipa-61d;
ORF Name	NTID	AAID	NT AA score probability
A17503000987_972187_£2_208	1672	5444	
Description gp:[GI:g2689564] [LN:U93688] [AC:U93 aureus] [DB:genpept-bct2] [DE:Staphy (tst),enterotoxin (ent), and integra staphylococcal phage integrase] [LE:	lococcus se (int)	s aure gene	us toxic shock syndrome toxin-1 s, complete cds.] [NT:similar to
ORF Name	NTID	AAID	NT AA score probability
A17503000987_9765677_c3_949	1673	5445	804 267 226 8.4e-19
Description gp:[GI:g928839] [LN:BK5TATTP] [AC:L4 phage BK5-T] [SR:Bacteriophage BK5-T ORF'410, 3' end pf cds, 20 ORFs, rep complete cds, ORF70'gene, 5' end of [DI:direct]	DNA] [I	OB:gen	pept-phg] [DE:Bacteriophage BK5-T , and Cro repressor protein genes,
ORF Name	NTID	AAID	NT AA score probability
A17503000987_977181_c1_702	1674	5446	<u> </u>
Description			
NO-HIT			
ORF Name	NTID	AAID	NT AA LengthLength score probability
AI7503000987_978377_c1_689	1675	5447	264 87 75 0.0037
Description			
<pre>sp:[LN:VG05_VACCC] [AC:P21026] [GN:G [DE:PROTEIN G5] [SP:P21026] [DB:swis protein] [OR:vaccinia virus] [DB:pir [OR:Vaccinia virus] [SR:Vaccinia vir [DB:genpept-vrl] [DE:Vaccinia virus, [RE:76522] [DI:direct]</pre>	sprot] > 2] >gp:{ us (stra	pir:[] [GI:g3] ain Co	LN:A42512] [AC:A42512] [PN:G5R 35417] [LN:VACCG] [AC:M35027] penhagen) DNA, clone VC-2]

ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	score	probability
AI7503000987_9819392_c3_908	1676	5448		262	179	6.4e-13
Description gp:[GI:g1463023] [LN:CELF45E1] [AC:U [SR:Caenorhabditis elegans strain=Br elegans cosmid F45E1.] [LE:13873:152 [DI:directJoin]	istol N2] [DB	genpep:	t-inv1] [DE:	Caenorhabditis
ORF Name A17503000987_9924055_c2_765 Description NO-HIT		<u>AAID</u> 5449	NT Length	AA Length 71	score	probability
ORF Name A17503000987 9926903 c2 835		<u>AAID</u>	<u>NT</u> Length]	AA Length	score	probability
Description pir: [LN:F45681] [AC:F45681] [PN:orf [LN:S57515] [AC:S57515] [GN:orf 61.2 {intergenic region between 41 and 61 sequence comes from Fig. A3.] [LE:12	61.2] [6] [OR:co] [bacte:	OR:phag liphag riopha	age T2] ge T2] age T2,	[DB:p: [DB:gen] [DB:genomic	ir2] >	gp:[GI:g298525] phg] [DE:orf 61.2
ORF Name A17503000987_994052_c3_979 Description gp:[GI:g4049717] [LN:AF063866] [AC:A [GN:MSV156] [OR:Melanoplus sanguinips sanguinipes entomopoxvirus, complete	1679 [F063866]	5451 [PN:0	DRF MSV	511 156 hyp DB:genp	pothet pept-v	rl] [DE:Melanoplus
ORF Name A17503000988_22939705_c3_35 Description		AAID 5452	NT Lengthl			probability
gp:[GI:g2689564] [LN:U93688] [AC:U93688] [DE:Staphy: (tst), enterotoxin (ent), and integras staphylococcal phage integrase] [LE:	lococcus se (int)	aurei	s, comp	c shock lete co	c synd ds.] [rome toxin-1
ORF Name			<u>NT</u> LengthI	Length.	score	probability
A17503000988_235762_f2_11 Description	1681	5453	159	52		
NO-HIT						
ORF Name	NTID 2	AAID	<u>NT</u> Length I	<u>AA</u> Length	score	probability
AI7503000988_23626577_c2_32	1682	5454	210	69	56	0.0035
Description gp:[GI:d1009788:g829068] [LN:SHFORF] [SR:Shigella sonnei (individual_isola [DE:Shigella sonnei DNA for 26 ORFs,	ate HW383	3) DNA	, clone	e pJK11	L42] [DB:genpept-bct1]

ORF Name	NTID	AAID LengthLength score probability
AI7503000988_24297062_f1_1	1683	5455 183 60
Description		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
AI7503000988_24642202_c1_30	1684	5456 648 215 822 5.8e-82
Description		
[DE:Staphylococcus aureus toxic shoc	k syndro	OR:Staphylococcus aureus] [DB:genpept-bct2] Tome toxin-1 (tst), enterotoxin (ent), and Orf7] [LE:6109] [RE:6708] [DI:complement]
ORF Name	NTID	AAID NT AA score probability
A17503000988_25665687_c2_33	1685	5457 876 291 322 5.6e-29
Description		
[DE:Staphylococcus aureus toxic shoc	k syndro	OR:Staphylococcus aureus] [DB:genpept-bct2] Come toxin-1 (tst),enterotoxin (ent), and Orf13] [LE:9717] [RE:10004] [DI:complement]
ORF Name	NTID	AAID NT AA score probability
AI7503000988_34428905_f1_3	1686	5458 165 54
Description		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000988_35428187_c3_37	1687	5459 438 145
Description	_	
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
	MIID	Length Length Diosability
AI7503000988_36601678_c1_29	1688	Length Length 50010 P10000011117
Description	1688	EngthLength 5460 1728 575 102 0.010
Description gp:[GI:g1004289] [LN:PPRNAE14B] [AC:	1688 Z50050]	[PN:Xylanase B] [OR:Penicillium
Description gp:[GI:g1004289] [LN:PPRNAE14B] [AC:	250050] 3.2.1.8]	[PN:Xylanase B] [OR:Penicillium] [DE:P.purpurogenum mRNA for
Description gp:[GI:g1004289] [LN:PPRNAE14B] [AC:purporogenum] [DB:genpept-pln1] [EC:	250050] 3.2.1.8]	[PN:Xylanase B] [OR:Penicillium] [DE:P.purpurogenum mRNA for DI:direct]
Description gp:[GI:g1004289] [LN:PPRNAE14B] [AC:purporogenum] [DB:genpept-pln1] [EC:	250050] 3.2.1.8]	[PN:Xylanase B] [OR:Penicillium] [DE:P.purpurogenum mRNA for
Description gp:[GI:g1004289] [LN:PPRNAE14B] [AC: purporogenum] [DB:genpept-pln1] [EC: endo-1,4-beta-xylanase.] [LE:11] [RE	Z50050] 3.2.1.8] :637] [I	[PN:Xylanase B] [OR:Penicillium [DE:P.purpurogenum mRNA for DI:direct] AAID NT AA score probability
Description gp:[GI:g1004289] [LN:PPRNAE14B] [AC: purporogenum] [DB:genpept-pln1] [EC: endo-1,4-beta-xylanase.] [LE:11] [RE ORF Name	Z50050] 3.2.1.8] :637] [I	[PN:Xylanase B] [OR:Penicillium] [DE:P.purpurogenum mRNA for DI:direct] AAID NT AA LengthLength score probability

ORF Name	NTID	AAID	NT AA score probability
A17503000988_9923176_£3_20	1690	5462	
Description			
elegans] [EC:1.6.5.3] [DB:pir2] >gp:	CHAIN 5, [PN:NA piquinon :[GI:g51 egans] [al genom] [SP: DH deh e) cha 5886] SR:Cae e.] [N	P24896] [DB:swissprot] ydrogenase (ubiquinone), chain 5] in 5] [OR:mitochondrion Caenorhabditis [LN:MTCE] [AC:X54252:S93745] norhabditis elegans] [DB:genpept-inv1]
ORF Name	NTID	AAID	NT AA LengthLength score probability
AI7503000990_10003756_£3_471	1691	5463	138 45
Description			
NO-HIT	of Management 27 To the Addition Action	- M. 0 10 10 10 10 10 10 10	
ORF Name	NTID	AAID	NT AA score probability
AI7503000990_10193763_c2_864	1692	5464	186 61
Description			
NO-HIT			
ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000990_10194713_f1_31	1693	5465	147 48
Description			
NO-HIT			
ORF Name	NTID	AAID	NT AA score probability
A17503000990_10241287_c2_822	1694	5466	663 220 74 0.037
Description		_	
<pre>gp:[GI:g924349] [LN:HIV1U13473] [AC: [GN:env] [OR:Human immunodeficiency immunodeficiency virus type 1 isolat (env) gene, V1V2 region, partial cds</pre>	virus to	ype 1] lone 0:	[DB:genpept-vrl] [DE:Human 3 fromUganda, envelope glycoprotein
ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000990_10268812_c2_820	1695	5467	144 47
Description			
NO-HIT			
ORF Name	NTID	AAID	NT AA score probability
AI7503000990_10360902_£3_536	1696	5468	183 60
Description	<u> </u>		d I
NO-HIT			

sp:[LN:POTD ECOLI] [AC:P23861] [GN:POTD] [OR:ESCHERICHIA COLI] [DE:SPERMIDINE/PUTRESCINE-BINDING PERIPLASMIC PROTEIN PRECURSOR (SPBP)] [SP:P23861] [DB:swissprot] >pir:[LN:D40840] [AC:D40840:H64856] [PN:spermidine/putrescine-binding protein precursor:spermidine/putrescine transport protein D] [GN:potD] [OR:Escherichia coli] [DB:pir2] >gp:[GI:d1036929:g1651550] [LN:D90747] [AC:D90747:AB001340] [PN:Spermidine/putrescine transport protein D] [GN:potD] [OR:Escherichia coli] [SR:Escherichia coli(strain:K12) DNA, clone:Kohara clone #238] [DB:genpept-bct1] [DE:Escherichia coli genomic DNA. (25.2 - 25.6 min).] [NT:ORF_ID:o238#12; similar to PIR Accession Number] [LE:13470] [RE:14516] [DI:complement] >qp:[GI:q1787367] [LN:AE000212] [AC:AE000212:U00096] [PN:spermidine/putrescine periplasmic transport] [GN:potD] [FN:transport; Transport of small molecules: Amino] [OR:Escherichia coli] [DB:genpept-bct2] [DE:Escherichia coli K-12 MG1655 section 102 of 400 of the completegenome.] [NT:f348; 100 pct identical to POTD ECOLI SW: P23861] [LE:7785] [RE:8831] [DI:complement] >gp:[GI:g147329] [LN:ECOPOTABCD] [AC:M64519] [PN:transport protein] [GN:potD] [OR:Escherichia coli] [SR:E.coli (strain DR112) DNA, clone pPT104] [DB:genpept-bct2] [DE:E.coli transport protein (potA, potB, potC and potD) genes, complete cds.] [LE:3144] [RE:4190] [DI:direct]

 ORF Name
 NTID
 AAID
 NT AA LengthLength
 Score
 probability

 A17503000990_10579000_c1_734
 1698
 5470
 957
 318
 728
 5.3e-72

Description

sp:[LN:COXX_BACSU] [AC:P24009] [GN:CTAB] [OR:BACILLUS SUBTILIS] [DE:PROBABLE CYTOCHROME C OXIDASE ASSEMBLY FACTOR] [SP:P24009] [DB:swissprot] >pir:[LN:C69609] [AC:C69609:S14395] [PN:cytochrome caa3 oxidase (assembly factor) ctaB] [GN:ctaB] [CL:heme O synthase] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e334763:g2339991] [LN:BS16823KB] [AC:Z98682] [PN:CtaB protein] [GN:ctaB] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis genomic DNA 23.9kB fragment.] [SP:P24009] [LE:4441] [RE:5358] [DI:direct] >gp:[GI:g994794] [LN:BSCTABF] [AC:X54140] [PN:cytochrome a assembly facto] [GN:ctaB] [OR:Bacillus subtilis] [DB:genpept-bct1] [EC:1.9.3.1] [DE:B. subtilis ctaB-F genes for cytochrome a assembly factor and cytochrome-c oxidase (EC 1.9.3.1) subunits II, I, II, and IVB.] [NT:putative] [SP:P24009] [LE:408] [RE:1325] [DI:direct] >gp:[GI:e1185078:g2633859] [LN:BSUB0008] [AC:Z99111:AL009126] [PN:cytochrome caa3 oxydase assembly factor] [GN:ctaB] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 8 of 21): from 1394791to 1603020.] [SP:P24009] [LE:163924] [RE:164841] [DI:direct]

 ORF Name
 NTID
 AAID
 NT Length Length
 Score
 probability

 A17503000990_1063753_c2_874
 1699
 5471
 1374
 457
 2070
 3.3e-214

Description

sp:[LN:MURD_STAAU] [AC:033595:007323] [GN:MURD] [OR:STAPHYLOCOCCUS AUREUS] [EC:6.3.2.9]
[DE:ADDING ENZYME)] [SP:033595:007323] [DB:swissprot] >pir:[LN:JC6560] [AC:JC6560]
[PN:UDP-N-acetylmuramoylalanine--D-glutamate ligase,

:UDP-N-acetylmuramoyl-L-alanyl-D-glutamate synthetase] [GN:murD] [OR:Staphylococcus aureus] [EC:6.3.2.9] [DB:pir2] >gp:[GI:g2271510] [LN:AF009671] [AC:AF009671] [PN:UDP-N-acetylmuramoyl-L-alanine : D-glutamate] [GN:murD] [FN:catalyzes ATP-dependent D-glutamate addition] [OR:Staphylococcus aureus] [DB:genpept-bct2] [DE:Staphylococcus aureus UDP-N-acetylmuramoyl-L-alanine : D-glutamateligase (murD) gene, complete cds.] [NT:MurD] [LE:1] [RE:1350] [DI:direct]

ORF Name AI7503000990_10740628_t3_637 Description NO-HIT	NTID 1700	AAID NT AA score probability 5472 183 60
ORF Name A17503000990_10928_c3_958 Description pir:[LN:T03492] [AC:T03492] [PN:hypotential of the properties of the pr	[LN:AFO	AAID NT AA LengthLength score probability 5473 1242 413 239 1.1e-17 al protein] [OR:Rhodobacter capsulatus] 10496] [AC:AF010496] [PN:hypothetical apept-bct2] [DE:Rhodobacter capsulatus
[LN:SAPTSHI] [AC:X93205] [PN:phosphoe	NTID 1702 SI] [OR: ME I)] enolpyrupt-bct1]	AAID NT AA LengthLength score probability 5474 1725 574 2602 1.4e-270 STAPHYLOCOCCUS AUREUS] [EC:2.7.3.9] [SP:P51183] [DB:swissprot] >gp:[GI:g1070386] avate-protein phosphatase] [GN:ptsI] [EC:2.7.3.9] [DE:S.aureus ptsH and ptsI
ORF Name AI7503000990_10972150_c3_1037 Description sp:[LN:RECG_STAAU] [AC:O50581] [GN:RI [DE:ATP-DEPENDENT DNA HELICASE RECG,] >gp:[GI:d1025491:g2826896] [LN:AB0004 [OR:Staphylococcus aureus] [SR:Staphylococcus [DB:genpept-bct1] [DE:Staphylococcus [RE:2379] [DI:direct]	[SP:05 439] [AC ylococci	50581] [DB:swissprot] C:AB000439] [PN:RecG] [GN:recG] us aureus (strain:RN4220) DNA]
PROTEIN IN LYTB-DAPB INTERGENIC REGIO [AC:JE0404:S40553:F64723:S22291] [PN protein] [OR:Escherichia coli] [EC:3. [AC:X54945] [GN:ORF 3] [OR:Escherichia interval.] [NT:product appears to be [DI:direct] >gp:[GI:d1001780:g216457] [AC:D10483:J01597:J01683:J01706:K0129 coli] [SR:Escherichia coli (strain:K- 0-2.4min. region.] [NT:hypothetical 3 [DI:direct] >gp:[GI:g1786213] [LN:AEG protein] [GN:yaaF] [FN:orf; Unknown]	ON] [SP: N:probak .2] ia coli] membrar [LN:EC 98:K0199 -12) DNF 32.6K pr 000113] [OR:Esc	O:M10420:M10611:M12544] [OR:Escherichia DB:genpept-bct1] [DE:E.coli K12 genome, otein(PIR:JE0404)] [LE:26947] [RE:27861] [AC:AE000113:U00096] [PN:orf, hypothetical cherichia coli] [DB:genpept-bct2] of 400 of the completegenome.] [NT:o304; 100

ORF Name			NT AA
Manue	NTID	AAID	LengthLength score probability
A17503000990_1178828_c2_810	1705	5477	705 234 649 1.3e-63
Description			
gp:[GI:g4097530] [LN:LLU64311] [AC:U [PN:phosphoribosylaminoimidazolesucc [DB:genpept-bct2] [EC:6.3.2.6] [DE:L phosphoribosylaminoimidazolesuccinoc phosphoribosylformylglycinamidine sy synthetase II (purL), andphosphoribo completecds; and unknown gene.] [NT:	inocarb actococ arboxam nthetas sylpyro	cus lac idesynt e I(pur phospha	thetase (purC), Q), phosphoribosylformylglycinamidine te amidotransferase (purF) genes,
ORF Name	NTID	AAID	NT AA LengthLength score probability
AI7503000990_1179542_c2_839	1706	5478	222 73
Description			
NO-HIT			
ORF Name	NTID	AAID	NT AA LengthLength
AI7503000990_11806512_£3_562	1707	5479	207 68 86 0.0050
Description			
gp:[GI:e1344613:g3874900] [LN:CEC41G elegans] [DB:genpept-inv1] [DE:Caeno [LE:18951:19702:20184] [RE:19549:200	rhabdit	is eleg	ans cosmid C41G6, complete sequence.]
ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000990_1204567_c2_897	1708	5480	750 249 481 8.0e-46
Description			
<pre>pir:[LN:G69878] [AC:G69878] [PN:con [CL:conserved hypothetical protein y [OR:Bacillus subtilis] [DB:pir2] >gp [AC:Z99112:AL009126] [GN:yloO] [FN:u</pre>	lo0:con :[GI:e1	served 185167:	hypothetical protein yloO homology]
[DE:Bacillus subtilis complete genom [NT:similar to hypothetical proteins >gp:[GI:e323505:g2337805] [LN:BSY139 [GN:yloO] [FN:unknown] [OR:Bacillus genomic DNA from the spoVM region.]] [LE:5 37] [AC subtili	ion 9 o 1363] [:Y13937 s] [DB:	cillus subtilis] [DB:genpept-bct1] f 21): from 1598421to 1807200.] RE:52127] [DI:direct]] [PN:putative Ptc1 protein] genpept-bct1] [DE:Bacillus subtilis
[DE:Bacillus subtilis complete genom [NT:similar to hypothetical proteins >gp:[GI:e323505:g2337805] [LN:BSY139 [GN:yloO] [FN:unknown] [OR:Bacillus] [LE:5 37] [AC subtili	ion 9 o 1363] [:Y13937 s] [DB: 35] [RE	cillus subtilis] [DB:genpept-bct1] f 21): from 1598421to 1807200.] RE:52127] [DI:direct]] [PN:putative Ptc1 protein] genpept-bct1] [DE:Bacillus subtilis
[DE:Bacillus subtilis complete genom [NT:similar to hypothetical proteins >gp:[GI:e323505:g2337805] [LN:BSY139 [GN:yloO] [FN:unknown] [OR:Bacillus genomic DNA from the spoVM region.]] [LE:5 37] [AC subtili [LE:131	ion 9 o 1363] [:Y13937 s] [DB: 35] [RE	cillus subtilis] [DB:genpept-bct1] f 21): from 1598421to 1807200.] RE:52127] [DI:direct]] [PN:putative Ptc1 protein] genpept-bct1] [DE:Bacillus subtilis :13899] [DI:direct] NT AA score probability
[DE:Bacillus subtilis complete genom [NT:similar to hypothetical proteins >gp:[GI:e323505:g2337805] [LN:BSY139 [GN:yloO] [FN:unknown] [OR:Bacillus genomic DNA from the spoVM region.] ORF Name	[LE:5 37] [AC subtili [LE:131] NTID	ion 9 o 1363] [:Y13937 s] [DB: 35] [RE AAID]	cillus subtilis] [DB:genpept-bct1] f 21): from 1598421to 1807200.] RE:52127] [DI:direct]] [PN:putative Ptc1 protein] genpept-bct1] [DE:Bacillus subtilis :13899] [DI:direct] NT AA LengthLength score probability [540] 179 [642] [6.9e-63]

ORF Name NTID AAID NT AA score probability
A17503000990_1209682_c2_856
Description
sp:[LN:SYFB_BACSU] [AC:P17922:P94540] [GN:PHET] [OR:BACILLUS SUBTILIS] [EC:6.1.1.20] [DE:TRNA LIGASE BETA CHAIN) (PHERS)] [SP:P17922:P94540] [DB:swissprot] >pir:[LN:YFBSB] [AC:A69676:I40460:S11731] [PN:phenylalaninetRNA ligase, beta chain:phenylalanyl-tRNA synthetase beta chain] [GN:pheT] [CL:phenylalaninetRNA ligase beta chain] [OR:Bacillus subtilis] [EC:6.1.1.20] [DB:pirl] >gp:[GI:e1184112:g2635328] [LN:BSUB0015] [AC:Z99118:AL009126] [PN:phenylalanyl-tRNA synthetase (beta subunit)] [GN:pheT] [OR:Bacillus subtilis] [DB:genpept-bct1] [EC:6.1.1.20] [DE:Bacillus subtilis complete genome (section 15 of 21): from 2795131to 3013540.] [SP:P17922] [LE:130980] [RE:133394] [DI:complement] >gp:[GI:e1165325:g1770031] [LN:BSZ75208] [AC:Z75208] [PN:phenylalanyl-tRNA synthetase beta subunit] [GN:pheT] [OR:Bacillus subtilis] [DB:genpept-bct1] [EC:6.1.1.20] [DE:B.subtilis genomic sequence 89009bp.] [NT:phenylalanyl-tRNA synthetase beta subunit] [SP:P17922] [LE:36513] [RE:38927] [DI:direct]
ORF Name NTID AAID NT AA score probability
A17503000990_13720312_c3_1024
Description
pir:[LN:C69878] [AC:C69878] [PN:hypothetical protein yloH] [GN:yloH] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1185160:g2633941] [LN:BSUB0009] [AC:Z99112:AL009126] [GN:yloH] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 9 of 21): from 1598421to 1807200.] [LE:43546] [RE:43749] [DI:direct] >gp:[GI:e323522:g2337798] [LN:BSY13937] [AC:Y13937] [PN:putative rpoZ protein] [GN:yloH] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis genomic DNA from the spoVM region.] [LE:5318] [RE:5521] [DI:direct]
ORF Name NTID AAID NT AA score probability
AI7503000990_13757785_f3_639
Description
NO-HIT
NTT 2.2
$rac{ ext{NTID}}{ ext{NTID}} = rac{ ext{NT}}{ ext{LengthLength}} rac{ ext{AA}}{ ext{score}} = ext{probability}$
A17503000990_1376317_c2_881
<u>Description</u>
gp:[GI:e199384:g1514599] [LN:LPPYRBSOP] [AC:Z54240] [GN:pyrR] [FN:regulatory protein] [OR:Lactobacillus plantarum] [DB:genpept-bct1] [DE:L.plantarum pyrimidine biosynthetic operon (pyrR, pyrB, pyrC,pyrAA, pyrAB, pyrD, pyrF, and pyrE) genes.] [LE:781] [RE:1323] [DI:direct]
ORF Name NTID AAID NT AA score probability
AI7503000990 13797076 c3 915 [1714] [5486 [135] [44

Description NO-HIT

ORF Name	NTID	AAID	NT AA LengthLength score probability
AI7503000990_13864213_£3_624	1715	5487	234 77 186 1.4e-14
subtilis] [DB:pir2] >gp:[GI:e118504 [GN:ykzG] [FN:unknown] [OR:Bacillus	4:g2633 s subtil	825] [I is] [DE	otein ykzG] [GN:ykzG] [OR:Bacillus LN:BSUB0008] [AC:Z99111:AL009126] B:genpept-bct1] [DE:Bacillus subtilis b 1603020.] [LE:129406] [RE:129615]
ORF Name	NTID	AAID	NT AA LengthLength probability
AI7503000990_13869091_£2_433	1716	5488	294 97 246 6.4e-21
[DE:SUBUNIT QOXD)] [SP:P34959] [DB:] [PN:quinol oxidase aa3-600 chain qoxD] [GN:qoxD] [OR:Bacillus subti [AC:M86548] [PN:quinol oxidase] [GN DNA] [DB:genpept-bct1] [DE:Bacillus QOXD)genes, complete cds.] [LE:4425 [AC:X73124] [GN:ipa-40d qoxD] [OR:E genomic region (325 to 333).] [SP:E >gp:[GI:e1186313:g2636349] [LN:BSUE oxidase (subunit IV)] [GN:qoxD] [OR	swisspronum (IV qoxD) Lis] [D] SigoxD] SigoxD[Sigox] [RE:4 Bacillus P34959] B0020] [RE:Bacillus P34959] [RE:Bacillus	ot] >pi :cytoch B:pir2] [OR:Bac is AA3- 799] [D subtil [LE:428 AC:Z991 us subt): from	cillus subtilis] [SR:Bacillus subtilis 600 quinol oxidase (QOXA, QOXB, QOXC, DI:direct] >gp:[GI:g413964] [LN:BSGENR] .is] [DB:genpept-bct1] [DE:B.subtilis 377] [RE:43251] [DI:direct] .23:AL009126] [PN:cytochrome aa3 quinol cilis] [DB:genpept-bct1] [DE:Bacillus a 3798401to 4010550.] [NT:alternate
ORF Name	NTID	AAID	<u>NT AA</u> LengthLength
AI7503000990_13916017_c1_738	1717	5489	123 40
Description			
NO-HIT			
ORF Name	NTID	AAID	NT AA score probability
A17503000990_14460882_c2_840	1718	5490	240 79 321 7.2e-29
Description gp:[GI:g1022726] [LN:SHU35635] [AC: haemolyticus] [SR:Staphylococcus ha [DE:Staphylococcus haemolyticus IS1 [LE:1101] [RE:1922] [DI:complement]	emolytic 272 ORF:	cus str	rain=Y176] [DB:genpept-bct1]
ORF Name	NTID	AAID	NT AA LengthLength score probability
AI7503000990_14557882_f3_532	1719	5491	

Description NO-HIT

NT ORF Name NTID probability LengthLength A17503000990 14642135 c3 957 1905 1720 4.1e-255 Description sp:[LN:TYPA BACSU] [AC:007631] [GN:TYPA] [OR:BACILLUS SUBTILIS] [DE:GTP-BINDING PROTEIN TYPA/BIPA HOMOLOG] [SP:007631] [DB:swissprot] >pir:[LN:E69872] [AC:E69872] [PN:GTP-binding elongation factor homolog ylaG] [GN:ylaG] [CL:translation elongation factor Tu homology] [OR:Bacillus subtilis] [DB:pir2] >qp:[GI:e1185067:q2633848] [LN:BSUB0008] [AC:Z99111:AL009126] [GN:ylaG] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 8 of 21): from 1394791to 1603020.] [NT:similar to GTP-binding elongation factor] [SP:007631] [LE:150736] [RE:152574] [DI:direct] >gp:[GI:e1191893:g2224766] [LN:BSZ97025] [AC:Z97025] [GN:ylaG] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis nprE, yla[A,B,C,D,E,F,G,H,I,J,K,L,M,N,O] and pycAgenes.] [NT:product highly similar to elongation factor EF-G] [SP:007631] [LE:4995] [RE:6833] [DI:direct] NT ORF Name NTID AAID score probability LengthLength AI7503000990 14644066 c2 865 1721 189 0.0032 Description gp:[GI:g3212079] [LN:AF068633] [AC:AF068633] [PN:phenol soluble modulin beta 1] [FN:inflammatory protein] [OR:Staphylococcus epidermidis] [DB:genpept-bct2] [DE:Staphylococcus epidermidis phenol soluble modulin beta 1 and phenolsoluble modulin beta 2 genes, complete cds.] [NT:PSM beta 1] [LE:669] [RE:803] [DI:direct] >gp:[GI:g3212080] [LN:AF068633] [AC:AF068633] [PN:phenol soluble modulin beta 2] [FN:inflammatory protein] [OR:Staphylococcus epidermidis] [DB:genpept-bct2] [DE:Staphylococcus epidermidis phenol soluble modulin beta 1 and phenolsoluble modulin beta 2 genes, complete cds.] [NT:PSM beta 2] [LE:859] [RE:993] [DI:direct] AΑ ORF Name NTID score probability LengthLength A17503000990 14650302_c1_688 1722 5494 264 8.0e-14 Description sp:[LN:YEXA BACSU] [AC:P12049] [GN:YEXA] [OR:BACILLUS SUBTILIS] [DE:HYPOTHETICAL 9.7 KD PROTEIN IN PURC-PURL INTERGENIC REGION] [SP:P12049] [DB:swissprot] >pir:[LN:E29326] [AC:E29326:E69799] [PN:conserved hypothetical protein yexA:hypothetical protein (pur operon)] [GN:yexA] [CL:conserved hypothetical protein MJ1593] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1182626:g2632960] [LN:BSUB0004] [AC:Z99107:AL009126] [GN:yexA] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 4 of 21): from 600701 to813890.] [NT:similar to hypothetical proteins] [LE:101124] [RE:101378] [DI:direct] NT AA NTID AAID

ORF Name score probability LengthLength AI7503000990 14664012 f2 323 1723 5495 468 155 341 5.4e-31

Description

pir:[LN:E69875] [AC:E69875] [PN:hypothetical protein ylbP] [GN:ylbP] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e334785:g2340013] [LN:BS16823KB] [AC:Z98682] [PN:Ylbp protein] [GN:ylbP] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis genomic DNA 23.9kB fragment.] [LE:21896] [RE:22378] [DI:complement] >gp:[GI:e1185100:g2633881] [LN:BSUB0008] [AC:Z99111:AL009126] [GN:ylbP] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 8 of 21): from 1394791to 1603020.] [LE:181379] [RE:181861] [DI:complement]

ORF Name	NTID	AAID	NT AA score	probability
A17503000990_14851587_c3_1010	1724	5496	126 41	
Description				
NO-HIT				
ORF Name	NTID	AAID	NT AA LengthLength	probability
A17503000990_14875002_f2_408	1725	5497		.0044
Description	L			
<pre>gp:[GI:g488889] [LN:A12521] [AC:A125 [OR:Plasmodium falciparum] [SR:malar [DE:Ag189 clone.] [LE:1:61] [RE:45:9</pre>	ia para	site P	. falciparum] [DB:ge	
ORF Name	NTID	AAID	NT AA LengthLength score	probability
A17503000990_156502_£1_179	1726	5498	243 80	
Description				
NO-HIT		and the same of the same of		
ORF Name	NTID	AAID	NT AA LengthLength score	probability
A17503000990_16593800_£2_359	1727	5499	144 47	
Description				
NO-HIT				
ORF Name	NTID	AAID	NT AA LengthLength	probability
A17503000990_19537562_f2_429	1728	5500	198 65	
Description				
NO-HIT				
ORF Name	NTID	AAID	NT AA LengthLength	probability
AI7503000990_19537878_f1_177	1729	5501	159 52	
Description				
NO-HIT				
ORF Name	NTID	AAID	NT AA LengthLength	probability
A17503000990_19556326_f1_94	1730	5502	141 46	
Description NO-HIT				

NT ORF Name NTID AAID score probability Length Length A17503000990 19557752 cl 777 1731 5503 912 2.7e-84 Description sp:[LN:PYRB BACSU] [AC:P05654] [GN:PYRB] [OR:BACILLUS SUBTILIS] [EC:2.1.3.2] [DE:TRANSCARBAMYLASE) (ATCASE)] [SP:P05654] [DB:swissprot] >pir:[LN:OWBSAC] [AC:A25015:C39845:B69686] [PN:aspartate carbamoyltransferase, catalytic chain:aspartate transcarbamoylase] [GN:pyrB] [CL:ornithine carbamoyltransferase: aspartate/ornithine carbamoyltransferase homology] [OR:Bacillus subtilis] [EC:2.1.3.2] [DB:pirl] [MP:37 min] >gp:[GI:g143384] [LN:BACPYRB] [AC:M13128] [OR:Bacillus subtilis] [SR:B.subtilis (strain JH861) DNA, clone pLS201] [DB:genpept-bct1] [DE:B.subtilis pyrB gene encoding aspartate transcarbamoylase, completecds.] [NT:aspartate transcarbamoylase (EC 2.1.3.2)] [LE:96] [RE:1010] [DI:direct] >gp:[GI:e1185141:g2633922] [LN:BSUB0009] [AC:Z99112:AL009126] [PN:aspartate carbamoyltransferase] [GN:pyrB] [FN:pyrimidine biosynthesis] [OR:Bacillus subtilis] [DB:genpept-bct1] [EC:2.1.3.2] [DE:Bacillus subtilis complete genome (section 9 of 21): from 1598421to 1807200.] [SP:P05654] [LE:21455] [RE:22369] [DI:direct] >gp:[GI:q143387] [LN:BACPYROP] [AC:M59757] [PN:aspartate transcarbamylase] [GN:pyrB] [OR:Bacillus subtilis] [DB:genpept-bct2] [DE:Bacillus subtilis ORF1A (pyrR), putative membrane-bound uracilpermease (pyrP), aspartate transcarbamylase (pyrB), dihydroorotase(pyrC), glutaminase of carbamyl phosphate synthetase (pyrAA), carbamyl phosphate synthetase (pyrAB), dihydroorotase dehydroqenase(pyrD), OMP decarboxylase (pyrF), and OMP-PRPP transferase (pyrE)genes, complete cds; and unknown gene.] [LE:2859] [RE:3773] [DI:direct] NT AA ORF Name NTID AAID score probability LengthLength A17503000990 19565627 c2 879 5504 132 1732 43 Description NO-HIT NT AAORF Name NTID AAID score probability LengthLength AI7503000990 19609530 f1 155 1733 5505 147 48 Description NO-HIT NT AΑ ORF Name NTID AAID score probability LengthLength A17503000990 19804703 c2 811 5506 747 248 1734 696 1.3e-68 Description sp:[LN:PURQ_BACSU] [AC:P12041] [GN:PURQ] [OR:BACILLUS SUBTILIS] [EC:6.3.5.3] [DE:SYNTHASE I)] [SP:P12041] [DB:swissprot] >pir:[LN:SYBS1G] [AC:F29326:H69684] [PN:phosphoribosylformylglycinamidine synthase, component I] [GN:purQ] [CL:phosphoribosylformylglycinamidine synthase component I] [OR:Bacillus subtilis] [EC:6.3.5.3] [DB:pir1] [MP:18 min] >gp:[GI:g143368] [LN:BACPURF] [AC:J02732:K00047] [OR:Bacillus subtilis] [SR:B.subtilis (strain DE1 (prototroph DER. or W168)) DNA, clone pPZ] [DB:genpept-bct1] [DE:B.subtilis pur operon encoding purine biosynthesis enzymes, 12genes.] [NT:phosphoribosylformyl glycinamidine synthetase I] [LE:4393] [RE:5076] [DI:direct] >gp:[GI:e1182627:g2632961] [LN:BSUB0004] [AC:Z99107:AL009126]

[PN:phosphoribosylformylglycinamidine synthetase II] [GN:purL] [FN:purine biosynthesis] [OR:Bacillus subtilis] [DB:genpept-bct1] [EC:6.3.5.3] [DE:Bacillus subtilis complete genome (section 4 of 21): from 600701 to813890.] [SP:P12041] [LE:101375] [RE:102058]

[DI:direct]

ORF Name	NTID	AAID LengthLength score probability
A17503000990_19957802_c1_752	1735	5507 1896 631 2764 9.5e-288
[GN:uvrC] [FN:excision of ultraviole	t light- aureus	trxA and uvrC genes and partial mutS and
ORF Name A17503000990_20312515_f2_242 Description NO-HIT	NTID 1736	AAID <u>NT AA</u> score probability 5508 141 46
ORF Name A17503000990_20500055_f1_218 Description NO-HIT	NTID 1737	AAID LengthLength score probability 5509 132 43
<pre>subtilis] [DB:pir2] >gp:[GI:e1185102 [GN:yllA] [FN:unknown] [OR:Bacillus</pre>	othetica :g263388 subtilis	AAID NT AA LengthLength score probability 5510 1626 541 718 6.1e-71 1 protein ylla] [GN:ylla] [OR:Bacillus 3] [LN:BSUB0008] [AC:Z99111:AL009126] [DB:genpept-bct1] [DE:Bacillus subtilis 791to 1603020.] [LE:182988] [RE:184607]
ORF Name A17503000990_2067627_c2_849 Description NO-HIT	NTID 1739	AAID NT AA score probability 5511 129 42
ORF Name A17503000990_20890875_c2_801 Description NO-HIT		AAID NT AA score probability 5512 147 48
ORF Name A17503000990_209840_f1_44 Description NO-HIT		AAID NT AA score probability 5513 195 64

ORF Name	NTID	AAID	NT AA score probability
AI7503000990_20994212_£1_151	1742	5514	207 68 71 0.022
Description	J I————	J [
complete genome (section 8 of 21): : [DI:complement] >gp:[GI:e324391:g222 [OR:Bacillus subtilis] [DB:genpept-}	6:g26338 subtil: from 139 24765] bct1] [I	347] [I is] [DE 94791tc [LN:BS2 DE:Baci	LN:BSUB0008] [AC:Z99111:AL009126] B:genpept-bct1] [DE:Bacillus subtilis D 1603020.] [LE:150435] [RE:150623] Z97025] [AC:Z97025] [GN:ylaF]
ORF Name	NTID	AAID	NT AA score probability
AI7503000990_211677_c3_989	1743	5515	702 233 436 4.7e-41
Description Description	•	1	
[DE:Bacillus subtilis complete genor [NT:similar to hypothetical proteins >gp:[GI:e1182723:g2633057] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-1 5 of 21): from 802821 to1011250.] [N	p:[GI:e] unknown] me (sect s] [LE:2 0005] [A bct1] [I NT:simi] 1020926: subtili	1182712 [OR:E zion 4 203465] AC:Z991 DE:Baci lar to :g21167	2:g2633046] [LN:BSUB0004] Bacillus subtilis] [DB:genpept-bct1] of 21): from 600701 to813890.] [RE:204172] [DI:complement] 108:AL009126] [GN:yfnB] [FN:unknown] illus subtilis complete genome (section hypothetical proteins) [LE:1345] 760] [LN:D86418] [AC:D86418] [PN:YfnB] rain:AC327) DNA] [DB:genpept-bct1]
ORF Name	NTID	AAID	NT AA LengthLength score probability
AT7503000990_21648962_c3_930	1744	5516	1827 608 1793 7.4e-185
Description gp:[GI:g517205] [LN:SPU09352] [AC:U0 streptococcal] [OR:Streptococcus pyo 42 KD protein (ORF1) gene and 67 KDN complete cds.] [NT:ORF2] [LE:1734]	ogenes] Myosin-c	[DB:ge crossre	enpept-bctl] [DE:Streptococcus pyogenes eactive streptococcal antigen gene,
ORF Name	NTID	AAID	NT AA LengthLength score probability
17503000990_21656327_c1_761	1745	5517	936 311 1464 5.4e-150
Description	,		
m · [GT · G2149891] [I.N · SAU94706] [AC · I	1947061	[DM·un	known [CN-v]]Cl [OP-Staphylococcus

aureus] [DB:genpept-bct2] [DE:Staphylococcus aureus strain ATCC 8325-4 cell wall/cell divisiongene cluster, yllB, yllC, yllD, pbpA, mraY, murD, div1B, ftsA andftsZ genes, complete cds.] [LE:987] [RE:1922] [DI:direct]

NT ORF Name NTID probability A17503000990 21759653_f3_666 5518 609 202 Description sp:[LN:QOX3 BACSU] [AC:P34958] [GN:QOXC:IPA-39D] [OR:BACILLUS SUBTILIS] [EC:1.9.3.-] [DE:SUBUNIT QOXC)] [SP:P34958] [DB:swissprot] >pir:[LN:C38129] [AC:C38129:S39694:G69687] [PN:bo-type ubiquinol oxidase, chain III qoxC:cytochrome aa3 quinol oxidase (subunit III) qoxC] [GN:qoxC] [CL:cytochrome-c oxidase chain III] [OR:Bacillus subtilis] [EC:1.10.3.-] [DB:pir2] >gp:[GI:g143398] [LN:BACQOXA] [AC:M86548] [PN:quinol oxidase] [GN:QOXC] [OR:Bacillus subtilis] [SR:Bacillus subtilis DNA] [DB:genpept-bct1] [DE:Bacillus subtilis AA3-600 quinol oxidase (QOXA, QOXB, QOXC, QOXD)genes, complete cds.] [LE:3809] [RE:4423] [DI:direct] >qp:[GI:q413963] [LN:BSGENR] [AC:X73124] [GN:ipa-39d goxC] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:B.subtilis genomic region (325 to 333).] [SP:P34958] [LE:42261] [RE:42875] [DI:direct] >gp:[GI:e1186314:g2636350] [LN:BSUB0020] [AC:Z99123:AL009126] [PN:cytochrome aa3 quino1 oxidase (subunit III)] [GN:qoxC] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 20 of 21): from 3798401to 4010550.] [NT:alternate gene name: ipa-39d] [SP:P34958] [LE:115087] [RE:115701] [DI:complement] NTAΑ ORF Name NTID AAID <u>sco</u>re probability LengthLength A17503000990 21915911 f1 197 5519 4.0e-19 Description pir: [LN:G69858] [AC:G69858] [PN:hypothetical protein ykoC] [GN:ykoC] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1181521:g2632041] [LN:BSAJ2571] [AC:AJ002571] [PN:YkoC] [GN:ykoC] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis 168 56 kb DNA fragment between xlyA and ykoR.] [LE:40195] [RE:40959] [DI:complement] >gp:[GI:e1183341:g2633675] [LN:BSUB0007] [AC:Z99110:AL009126] [GN:ykoC] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 7 of 21): from 1194391to 1411140.] [LE:193144] [RE:193908] [DI:complement] NTAΑ ORF Name NTID AAID score probability LengthLength A17503000990 22000943 f1 216 5520 222 Description NO-HIT NT AA ORF Name NTID AAID score probability LengthLength AI7503000990 22400283 c2 830 5521 981 326 1749 1329 1.1e-135 Description pir: [LN:C36718] [AC:C36718:A69674] [PN:pyruvate dehydrogenase (lipoamide), El beta chain precursor pdhB] [GN:pdhB] [CL:pyruvate dehydrogenase (lipoamide) beta chain] [OR:Bacillus subtilis] [EC:1.2.4.1] [DB:pir2] >gp:[GI:g143378] [LN:BACPYDHY] [AC:M57435:M31542] [PN:pyruvate decarboxylase (E-1) beta subunit] [GN:pdhB] [OR:Bacillus subtilis] [SR:B.subtilis (strain 168) BRB1 (sacA321 metB5) cell line DNA, clone] [DB:genpept-bct1] [EC:1.2.4.1] [DE:B.subtilis pyruvate dehydrogenase complex genes, complete cds; PAL-related lipoprotein (slp) gene, complete cds, lysinedecarboxylase (cad) gene, partial cds.] [LE:2796] [RE:3773] [DI:direct] >gp:[GI:e1185049:g2633830] [LN:BSUB0008] [AC:Z99111:AL009126] [PN:pyruvate dehydrogenase (E1 beta subunit)] [GN:pdhB] [OR:Bacillus subtilis] [DB:genpept-bct1] [EC:1.2.4.1] [DE:Bacillus subtilis complete genome (section 8 of 21): from 1394791to 1603020.] [SP:P21882] [LE:134060] [RE:135037] [DI:direct] >gp:[GI:g3282143] [LN:AF012285] [AC:AF012285:AF012284:U51911] [PN:pyruvate decarboxylase E-1 beta

subunit] [GN:pdhB] [OR:Bacillus subtilis] [DB:genpept-bct2] [EC:1.2.4.1] [DE:Bacillus subtilis mobA-nprE gene region.] [NT:similar to pyruvate decarboxylase (E-1) beta]

[LE:34548] [RE:35525] [DI:direct]

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000990_2242938_c3_1009	1750	5522	933	310	1055	1.2e-106
Description						
sp:[LN:YLYB_BACSU] [AC:Q45480:O31732 33.7 KD PROTEIN IN LSP-PYRR INTERGEN [DB:swissprot] >gp:[GI:g1373157] [LN [DB:genpept-bct1] [DE:Bacillus subticds,isoleucyl-tRNA synthetase (iles) hypothetical protein; Method: concep	IC REGI :BSU488 lis sig and py	ON (OR 370] [A gnal pe yrR gen	F-X)] C:U488' ptidase es, pa:	[SP:Q45 70] [OF e II (] rtial c	6480:0 R:Baci .sp) g :ds.]	31732] llus subtilis] ene, complete [NT:orf-X;
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000990_22461078_c1_769	1751	5523	735	244	814	4.1e-81
Description gp:[GI:g1314301] [LN:SAU41072] [AC:U aureus] [DB:genpept-bct2] [DE:Staphy gene,partial cds.] [LE:106] [RE:723]	lococci	ıs aure				
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
AI7503000990_22462802_c3_1026	1752	5524	1203	400	1070	3.1e-108
Description						
[DB:pir2] >gp:[GI:e1185161:g2633942] [FN:unknown] [OR:Bacillus subtilis] genome (section 9 of 21): from 15984 metabolism flavoprotein] [LE:43830] [LN:BSY13937] [AC:Y13937] [PN:putati subtilis] [DB:genpept-bct1] [DE:Baci [LE:5602] [RE:6822] [DI:direct]	[DB:ger 21to 18 [RE:450 ve Dfp	npept-b 307200. 50] [D protei	ct1] [I [NT:s :direc [GN:	DE:Baci similar ct] >gp :yloI]	llus ; to po :[GI:	subtilis complete antothenate e323501:g2337799] nknown] [OR:Bacillus
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
AI7503000990_22478427_c3_987	1753	5525	138	45	115	4.8e-07
Description						
sp:[LN:GGI2_STAHA] [AC:P11698] [OR:S PROTEIN 2 (GONOCOCCAL GROWTH INHIBIT [AC:S00600] [PN:antibacterial prote [CL:Staphylococcus haemolyticus anti [DB:pir1]	OR 2)] in 2:gc	[SP:P1] nococca	1698] al grov	[DB:swi wth inh	sspro	t] >pir:[LN:BXSA2] r 2]
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000990_22542567_c2_855	1754	5526	759	252	562	2.1e-54
Description						
pir:[LN:G69984] [AC:G69984] [PN:rRN.subtilis] [DB:pir2] >gp:[GI:e1184114 [GN:ysgA] [FN:unknown] [OR:Bacillus complete genome (section 15 of 21): methylase] [LE:134799] [RE:135545] [LN:BSZ75208] [AC:Z75208] [PN:hypothe [DB:genpept-bct1] [DE:B.subtilis general complete general co	:g26353 subtili from 27	30] [Li s] [DB 95131to	1:BSUB(genper 30135	0015] [pt-bct1 540.] [AC:Z99] [DE NT:sir	9118:AL009126] :Bacillus subtilis milar to rRNA

NT ORF Name NTID score probability LengthLength AI7503000990 22775126 ±2 411 1755 5527 633 210 1.0e-22 263 Description pir:[LN:A69859] [AC:A69859] [PN:hypothetical protein ykoE] [GN:ykoE] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1181523:q2632043] [LN:BSAJ2571] [AC:AJ002571] [PN:YkoE] [GN:ykoE] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis 168 56 kb DNA fragment between xlyA and ykoR.] [LE:42565] [RE:43164] [DI:complement] >gp:[GI:e1183343:g2633677] [LN:BSUB0007] [AC:Z99110:AL009126] [GN:ykoE] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 7 of 21): from 1194391to 1411140.] [LE:195514] [RE:196113] [DI:complement] NTLengthLength score ORF Name NTID AAID probability A17503000990_22931642_c2_802 5528 12242.6e-127 Description gp:[GI:d1024918:g2696796] [LN:AB009635] [AC:AB009635] [PN:Fmt] [GN:fmt] [OR:Staphylococcus aureus] [SR:Staphylococcus aureus (strain:KSA8) DNA] [DB:genpept-bct1] [DE:Staphylococcus aureus DNA for Fmt, complete cds.] [LE:1234] [RE:2427] [DI:direct] NT AΑ LengthLength score ORF Name NTID AAID probability A17503000990 23438887 c1 748 300 1.3e-15 Description pir: [LN:A69985] [AC:A69985] [PN:hypothetical protein yshA] [GN:yshA] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1184110:g2635326] [LN:BSUB0015] [AC:Z99118:AL009126] [GN:yshA] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 15 of 21): from 2795131to 3013540.] [LE:129612] [RE:129869] [DI:complement] >qp:[GI:e1165327:q1770033] [LN:BSZ75208] [AC:Z75208] [PN:hypothetical protein] [GN:yshA] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:B.subtilis genomic sequence 89009bp.] [NT:unknown function; putative] [LE:40038] [RE:40295] [DI:direct] NT AA ORF Name NTID AAID score probability LengthLength A17503000990 23442177 F3 600 1758 669 Description pir: [LN:D69864] [AC:D69864] [PN:hypothetical protein yktB] [GN:yktB] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1185055:g2633836] [LN:BSUB0008] [AC:Z99111:AL009126] [GN:yktB] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 8 of 21): from 1394791to 1603020.] [LE:140850] [RE:141488]

[DI:complement] >gp:[GI:g3282149] [LN:AF012285] [AC:AF012285:AF012284:U51911] [PN:unknown] [GN:yktB] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct2]

[DE:Bacillus subtilis mobA-nprE gene region.] [LE:41338] [RE:41976] [DI:complement]

ORF Name	NTID	AAID	NT AA LengthLength score probabil	ity
AI7503000990_23448838_t1_217	1759	5531	873 290 786 3.8e-78	
Description sp:[LN:FOLD_BACSU] [AC:P54382] [GN:FildE:METHENYLTETRAHYDROFOLATE CYCLOHY. >pir:[LN:E69626] [AC:E69626] [PN:memethenyltetrahydrofolate cyclohydroldehydrogenase (NAD+): methylenetetral[OR:Bacillus subtilis] [EC:1.5.1.5:3] [LN:BACJH642] [AC:D84432:D82370] [PN (strain:JH642(trpC2 PheA1)) DNA] [DB region containing skin element.] [LE >gp:[GI:e1185699:g2634865] [LN:BSUBO [PN:methenyltetrahydrofolate cyclohybiosynthesis] [OR:Bacillus subtilis] [DE:Bacillus subtilis complete genome [NT:alternate gene name: yqiA;] [SP:	DROLASE, thylenet ase,] [6 hydrofol .5.4.9] :YqiA] [:genpept :190351] 013] [AC drolase] [DB:gen e (secti	cetrahy En:foli late de [DB:pa [OR:Bac -bct1] [RE:1 [GN:f] [GN:f] [GN:f]]	P54382] [DB:swissprot] drofolate dehydrogenase (NAD] [CL:methylenetetrahydrofo hydrogenase (NAD+) homology] r2] >gp:[GI:d1013251:g130391 illus subtilis] [SR:Bacillus [DE:Bacillus subtilis DNA, 91202] [DI:direct] 6:AL009126] olD] [FN:purines and amino a ct1] [EC:1.5.1.5:3.5.4.9] of 21): from 2395261to 26137	P+), / late 6] subtilis 283 Kb cids 30.]
ORF Name A17503000990_23453767_f1_62 Description NO-HIT	NTID 1760	<u>AAID</u> 5532	NT AA score probabil LengthLength 138 45	ity
ORF Name AI7503000990_23492327_f1_37 Description pir: [LN:S75993] [AC:S75993] [PN:hypothetical [SR:Synechocystis sp. (strain:PCC6803 PCC6803 complete genome, 25/27, 31386 [RE:80528] [DI:complement]	B:pir2] l protei 3) DNA]	5533 al prot >gp:[G n] [GN [DB:ge	I:d1011491:g1001353] [LN:SYC :clpP] [OR:Synechocystis sp. npept-bct1] [DE:Synechocysti	[SR:PCC SLLLH]] s sp.
ORF Name AI7503000990_23532327_c2_829 Description sp:[LN:ODPA_BACST] [AC:P21873] [GN:PI [DE:PYRUVATE DEHYDROGENASE E1 COMPONI	DHA] [OR	5534 :BACII		1.2.4.1] prot]
A17503000990_23550313_f2_329 Description NO-HIT	1763	5535	141 46	
ORF Name AI7503000990_23572178_c2_861 Description sp:[LN:DHSB_BACSU] [AC:P08066] [GN:SI [DE:SUCCINATE DEHYDROGENASE IRON-SULE	1764 DHB] [OR	5536	NT AA score probabil: 846 281 1070 3.1e-108 LUS SUBTILIS] [EC:1.3.99.1] [SP:P08066] [DB:swissprot]	Lty

			2777	7. 7.		
ORF Name	NTID	AAID	NT Length	<u>AA</u> Lengtl	score	probability
A17503000990_23572253_c1_796	1765	5537	741	246	- 623	7.1e-61
Description	<u> </u>		J		-	
pir:[LN:B69693] [AC:B69693:JC4821]] [CL:ribonuclease III:double-strand subtilis] [EC:3.1.26.3] [DB:pir2] >g [AC:Z99112:AL009126] [PN:ribonuclease [DB:genpept-bct1] [EC:3.1.26.3] [DE:21): from 1598421to 1807200.] [NT:al [DI:direct]	ded RNA- gp:[GI:e se III] Bacillu	binding 1185184 [GN:rnd s subti	repea 1:g2633 cS] [OF	at homo 3965] R:Bacil omplete	ology] [LN:BSI llus si e genor	[OR:Bacillus UB0009] ubtilis] me (section 9 of
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Lengtl	score	probability
A17503000990_23632962_c2_880	1766	5538	810	269	270	1.8e-23
Description				L		
sp:[LN:YFIE_BACSU] [AC:P54721] [GN:YKD PROTEIN IN GLVBC 3'REGION] [SP:P5 [PN:conserved hypothetical protein y >gp:[GI:el182814:g2633148] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-b5 of 21): from 802821 to1011250.] [N subtilis] [SP:P54721] [LE:94696] [RE [LN:D50543] [AC:D50543] [PN:unknown] [SR:Bacillus subtilis (strain:168, h [DE:Bacillus subtilis DNA for 76-deg [DI:direct]	74721] [1 7fiE] [GI 7005] [A 7005] [A 7005] [A 7005] [DI 71:simila 71:95553] 81:95553]	DB:swis N:yfiE C:Z991(E:Bacil ar to h [DI:di iE] [FN e:haplo	ssprot]] [OR: 08:AL00 .lus su hypothe .rect] I:unkno	>pir Bacill 99126] ubtilis etical >gp:[G own] [G	:[LN:He lus sub [GN:yr s compl prote: prote: GI:d100 CR:Bac: B:genpe	[AC:H69802] [AC:H69802] otilis] [DB:pir2] fiE] [FN:unknown] lete genome (section ins from B. D9744:g1486247] fillus subtilis] ept-bct1]
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
AI7503000990_23642217_c1_790	1767	5539	648	215	396	8.1e-37
Description						
pir:[LN:C69879] [AC:C69879] [PN:hyp subtilis] [DB:pir2] >gp:[GI:el185171 [GN:yloS] [FN:unknown] [OR:Bacillus complete genome (section 9 of 21): f [DI:direct] >gp:[GI:e323508:g2337809 [GN:yloS] [FN:unknown] [OR:Bacillus genomic DNA from the spoVM region.]	g26339! subtilia rom 1598 [LN:BS subtilia	52] [LN s] [DB: 8421to SY13937 s] [DB:	:BSUB0 genpep 180720] [AC: genpep	009] bt-bctl 0.] [I Y13937 bt-bctl	[AC: Z99 L] [DE: LE: 5570 7] [PN: L] [DE:	9112:AL009126] :Bacillus subtilis 09] [RE:56353] :YloS protein] :Bacillus subtilis
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000990_23647178_c2_903	1768	5540		210	·	5.0e-37
Description		•				
<pre>pir:[LN:A69880] [AC:A69880] [PN:hyp subtilis] [DB:pir2] >gp:[GI:e1185179 [GN:ylpC] [FN:unknown] [OR:Bacillus complete genome (section 9 of 21): f</pre>	:g263396 subtilis	50] [LN s] [DB:	:BSUB0 genpep	009] t-bct1	[AC: Z99	9112:AL009126] Bacillus subtilis

[DI:direct] >gp:[GI:e323513:g2337817] [LN:BSY13937] [AC:Y13937] [PN:YlpC protein] [GN:ylpC] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis genomic DNA from the spoVM region.] [LE:24718] [RE:25284] [DI:direct]

ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	score	probability
AI7503000990_23650293_c1_785	1769	5541	627	208	693	2.7e-68
Description pir:[LN:B69878] [AC:B69878] [PN:guar [CL:guanylate kinase:guanylate kinase >gp:[GI:e1185159:g2633940] [LN:BSUB00 [OR:Bacillus subtilis] [DB:genpept-bo 9 of 21): from 1598421to 1807200.] [I [RE:43542] [DI:direct] >gp:[GI:e32350 Gmk protein] [GN:yloD] [FN:unknown] [DE:Bacillus subtilis genomic DNA fro [DI:direct]	e homolo 009] [A ct1] [D NT:simi 00:g233 [OR:Bac	ogy] [9 C:Z991 E:Baci lar to 7797] illus	OR:Baci 12:ALO llus su guanyl [LN:BSY subtili	illus s 09126] ubtilis Late ki (13937] [DB	ubtil: [GN:y] compliance nase] [AC:Y]	LoD] [DB:pir2] LoD] [FN:unknown] Lete genome (section [LE:42808] [213937] [PN:putative [2pt-bct1]
ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	score	probability
A17503000990_23695900_c2_812	1770	5542	1551	516		9.5e-153
gp:[GI:g4097534] [LN:LLU64311] [AC:U6 amidotransferase] [GN:purF] [OR:Lacto [DE:Lactococcus lactis phosphoribosy] phosphoribosylformylglycinamidine synsynthetase II (purL), andphosphoriboscompletecds; and unknown gene.] [NT:I	ococcus laminoin nthetase sylpyron PRPP ATa	lactianidazo e I(pum phospha ase] [1	s] [DB: lesucci rQ), ph ate ami	genpep nocarb nosphor dotran	ot-bct2 oxamic ibosyl	P] [EC:2.4.2.14] desynthetase (purC), formylglycinamidine se (purF) genes, [DI:direct]
ORF Name	NTID	AAID	<u>Length</u>		score	probability
A17503000990_23730340_c1_716 Description	1771	5543	1422	473	2270	2.1e-235
pir:[LN:S19723] [AC:S19723] [PN:dihy complex chain E3] [GN:pdhD] [CL:dihy dehydrogenase homology] [OR:Staphylocosyp:[GI:g48874] [LN:SAPDHDNA] [AC:X58 subunit E3] [GN:pdhD] [OR:Staphylocosy [DE:S.aureus pdhB, pdhC and pdhD generacetyltransferase and dihydrolipoamic	ydrolipo coccus a 8434:S73 ccus au es for p	oamide aureus] 3625] reus] oyruvat	dehydr [EC:1 [PN:dih [DB:gen ce deca	rogenas 8.1.4 nydroli npept-b nrboxyl	e:dihy] [DB: poamic ct1] ase,di	rdrolipoamide pir1] de dehydrogenase: [EC:1.8.1.4] .hydrolipoamide
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000990_23836036_c2_887	1772	5544		63		1.2e-09
Description sp:[LN:CARB_BACCL] [AC:P46537] [GN:PN] [DE:(EC 6.3.5.5) (CARBAMOYL-PHOSPHATE [DB:swissprot] >pir:[LN:I40169] [AC:I] (glutamine-hydrolyzing),] [GN:pyrAb] (glutamine-hydrolyzing) large chain:R synthase (glutamine-hydrolyzing) large 6.3.5.5] [DB:pir2] >gp:[GI:g312443] [Synthase] [GN:PyrAb] [OR:Bacillus cal [DE:B.caldolyticus pyrimidine biosynt [DI:direct]	E SYNTHI	ETASE A E34321 Arbamoy carboxy n homol (R] [AC cus] [I	AMMONIA] [PN: /l-phos /lase h logy] [C:X7330 DB:genp	CHAIN carbam phate comolog OR:Bac PN eept-bc)] [SF oyl-ph syntha y:carb illus :carba t1] [E	e:P46537] cosphate synthase cse camoyl-phosphate caldolyticus] [EC: cmoyl-phosphate cC:6.3.5.5]
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
AI7503000990_23928937_c3_953	1773	5545		46		
Description NO-HIT						

ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	score	probability
AI7503000990_24025253_±1_219	1774	5546	321	106	198	1.3e-14
Description pir:[LN:T00323] [AC:T00323] [PN:chi [EC:3.2.1.14] [DB:pir3] >gp:[GI:d102 [PN:chitinase B] [GN:chiB] [OR:Clost paraputrificum (strain:M21) DNA] [DB paraputrificum gene for chitinase B,	4701:g2 ridium :genpep	696017 parapu t-bct1] [LN:A trificu] [EC:3	B001874 m] [SR .2.1.14	4] [AC :Clost 4] [DE	:AB001874] ridium :Clostridium
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000990_24025467_c2_863	1775	5547	522	173	238	4.5e-20
Description sp:[LN:YSNB_BACSU] [AC:P94559] [GN:Y KD PROTEIN IN RPH-ILVB INTERGENIC RE [AC:D69986] [PN:conserved hypotheti hypothetical protein MG207: phosphoe [DB:pir2] >gp:[GI:e1184084:g2635300] [FN:unknown] [OR:Bacillus subtilis] genome (section 15 of 21): from 2795 proteins] [SP:P94559] [LE:103990] [R >gp:[GI:e1165358:g1770061] [LN:BSZ75 [GN:ysnB] [OR:Bacillus subtilis] [DB 89009bp.] [NT:homology to HI0260 of [RE:65917] [DI:direct]	GION] [cal pro sterase [LN:BS] [DB:gen] 131to 3 E:10450 208] [A	SP:P94! tein ys core l UB0015 pept-bo 013540 5] [DI C:Z7520 t-bct1]	559] [D snB] [G nomolog [AC:Z ct1] [D .] [NT: :comple [DB:B	B:swiss N:ysnB y] [OR 99118:2 E:Baci similar ment] :hypotl	sprot]] [CL :Bacil AL0091 llus s r to h hetica lis ge	>pir:[LN:D69986] :conserved lus subtilis] 26] [GN:ysnB] ubtilis complete ypothetical l protein] nomic sequence
ORF Name AI7503000990_24220002_f1_67 Description	NTID 1776	<u>AAID</u> 5548	NT Length]	AA Length	score	probability
NO-HIT				· . · . · . · . · . · . · . · . · . · .		
ORF Name	NTID	AAID	<u>NT</u> Lengthl	<u>AA</u> Length	score	probability
A17503000990_24240676_t3_477 Description NO-HIT	1777	5549	246	81		
ORF Name	NTID	AAID	<u>NT</u> Length I	<u>AA</u> Length	score	probability
A17503000990_24245437_c1_763 Description	1778	5550	1005		1471	9.8e-151
gp:[GI:d1023423:g2463562] [LN:AB0075 [OR:Staphylococcus aureus] [SR:Staphylococcus MraY,MurD, partial and complete cds. [LN:AF034153] [AC:AF034153] [PN:phos [OR:Staphylococcus aureus] [DB:genpe phospho-N-acetylmuramoyl-pentapeptid [RE:993] [DI:direct]	ylococci aureus] [LE:3: pho-N-ac pt-bct2	us aure genes 120] [F cetylmu] [DE:S	eus (st for per RE:4085 ramoyl- Staphylo	rain:NO nicilli [DI:d -pentar ococcus	CTC832 in-bindirect peptides aure	5) DNA] ding protein 1,] >gp:[GI:g4104230] e] [GN:mraY] us
ORF Name	NTID	AAID	<u>NT</u> LengthI	<u>AA</u> Length	score	probability
A17503000990_24256250_£3_467	1779	5551		41		
Description NO-HIT						

ORF Name	NTID	AAID	NT A	AA ngth score	probability
AI7503000990_24256551_c3_923	1780	5552	1071 35		3.0e-94
Description sp:[LN:PUR5_BACSU] [AC:P12043] [GN:P10E:(PHOSPHORIBOSYL-AMINOIMIDAZOLE STORM	YNTHETA H29326: e cyclo bosylfo clo-lig >gp:[GI s (stra s pur o idazole 4] [LN: thetase ct1] [E	SE) (A A69685 -ligase rmylgly ase hor :g1433' in DE1 peron e synthe BSUB000] [GN:p C:6.3.3	IR SYNTHAS	oribosylar ne cyclo-l DR:Bacillu ACPURF] [A Dph DER. c Durine bic R-M)] [LE: 99107:AL00 :purine bis Bacillus s	minoimidazole ligase: us subtilis] AC:J02732:K00047] or W168)) DNA, clone osynthesis enzymes, :8796] [RE:9836] 09126] iosynthesis] subtilis complete
ORF Name A17503000990_24257877_c2_854 Description NO-HIT	NTID 1781	<u>AAID</u> 5553	NT A Length Len	AA agth	probability
ORF Name AI7503000990 24261068 c1 758	NTID 1782	<u>AAID</u> 5554	NT A LengthLen	AA agth	probability
Description gp:[GI:g3212079] [LN:AF068633] [AC:AI [FN:inflammatory protein] [OR:Staphyl [DE:Staphylococcus epidermidis phenol beta 2 genes, complete cds.] [NT:PSM >gp:[GI:g3212080] [LN:AF068633] [AC:AI [FN:inflammatory protein] [OR:Staphyl [DE:Staphylococcus epidermidis phenol beta 2 genes, complete cds.] [NT:PSM	lococcus l solub beta 1 AF06863 lococcus l solub	s epide le modu] [LE:6 3] [PN: s epide le modu	ermidis] (alin beta 669] [RE:8 phenol sc ermidis] [alin beta	[DB:genper 1 and phe 303] [DI:doluble mod [DB:genper 1 and phe	enolsoluble modulin direct] dulin beta 2] bt-bct2]
ORF Name	NTID		<u>LengthLen</u>		probability
AI7503000990_24261068_c3_988 Description	1783	5555	147 48	215	1.2e-17
gp:[GI:g3212079] [LN:AF068633] [AC:AF [FN:inflammatory protein] [OR:Staphy] [DE:Staphylococcus epidermidis phenol beta 2 genes, complete cds.] [NT:PSM >gp:[GI:g3212080] [LN:AF068633] [AC:AF [FN:inflammatory protein] [OR:Staphy] [DE:Staphylococcus epidermidis phenol beta 2 genes, complete cds.] [NT:PSM	lococcus l solubl beta 1] AF068633 lococcus l solubl	s epide le modu [LE:6] [PN: s epide le modu	ermidis] [alin beta 669] [RE:8 phenol so ermidis] [alin beta	DB:genpep 1 and phe 03] [DI:d luble mod DB:genpep 1 and phe	ot-bct2] enolsoluble modulin direct] dulin beta 2] ot-bct2] enolsoluble modulin
ORF Name AI7503000990_24275137_f1_21 Description	NTID 1784	<u>AAID</u> 5556	NT A Length Len	- score	probability
NO-HIT					

ORF Name	NTID	AAID Length Length score probability
A17503000990_24297217_c1_780	1785	5557 3114 1037 3756 0.0
Description	اـــــا ك	
[DE: (EC 6.3.5.5) (CARBAMOYL-PHOSPHA [DB:swissprot] >pir: [LN:I40169] [AC (glutamine-hydrolyzing),] [GN:pyrAb (glutamine-hydrolyzing) large chain synthase (glutamine-hydrolyzing) la 6.3.5.5] [DB:pir2] >gp: [GI:g312443] synthase] [GN:PyrAb] [OR:Bacillus co	TE SYNTH C:140169: c) [CL:c c:biotin arge chai [LN:BCP caldolyti	S34321] [PN:carbamoyl-phosphate synthase carbamoyl-phosphate synthase carboxylase homology:carbamoyl-phosphate n homology] [OR:Bacillus caldolyticus] [EC:PYR] [AC:X73308] [PN:carbamoyl-phosphate
ORF Name	NTID	AAID NT AA score probability
A17503000990_24330337_f3_557	1786	5558 957 318 555 1.1e-53
Description		
pir:[LN:H69984] [AC:H69984] [PN:co	nserved	hypothetical protein ysgB] [GN:ysgB]
[OR:Bacillus subtilis] [DB:pir2] >g		
		[OR:Bacillus subtilis] [DB:genpept-bct1]
		ion 15 of 21): from 2795131to 3013540.]
[NT:similar to hypothetical protein		
>gp:[GI:e1165326:g1770032] [LN:BSZ7		
		t-bctl] [DE:B.subtilis genomic sequence LE:38963] [RE:39904] [DI:complement]
	acrec, [DE.30903] [KE.39904] [DI.Complement]
ORF Name	NTID	AAID NT AA score probability
		LengthLength Score probability
A17503000990_24351577_c1_677	1787	5559 171 56
Description		
NO-HIT		
		NITT 2.2
ORF Name	NTID	AAID LengthLength score probability
A17503000990_24355342_c1_799	1788	5560 1014 338 1265 6.6e-129
Description		
		:BACILLUS SUBTILIS] [DE:SIGNAL RECOGNITION
		P37105] [DB:swissprot] >pir:[LN:B47154]
		particle chain ffh] [GN:ffh] [CL:signal
recognition particle 54K protein] [
>gp:[G1:e1185189:g2633970] [LN:BSUB	0009] [A	C:Z99112:AL009126] [PN:signal recognition
		DB:genpept-bct1] [DE:Bacillus subtilis
		8421to 1807200.] [LE:73153] [RE:74493] D14356] [AC:D14356] [PN:Ffh] [GN:ffh]
		D14356] [AC:D14356] [PN:FIN] [GN:FIN] s (strain:168) DNA] [DB:genpept-bct1]
		for ORF1, Ffh and 30Sribosomal protein S16,
complete cds.] [LE:711] [RE:2051]		

ORF Name	NTID	AAID	NT LengthL	AA Length sc	ore	probability
A17503000990_24406291_c2_902	1789	5561			59 5	5.9e-34
Description	<u> </u>				IL	
pir:[LN:D69879] [AC:D69879] [PN:alk [OR:Bacillus subtilis] [DB:pir2] >gp [AC:Z99112:AL009126] [GN:yloU] [FN:u [DE:Bacillus subtilis complete genom [NT:similar to alkaline-shock protei >gp:[GI:e323527:g2337812] [LN:BSY139 [GN:yloU] [FN:unknown] [OR:Bacillus genomic DNA from the spoVM region.]	e:[GI:e1] nknown] e (sect: n] [LE:! 37] [AC subtilis	185174 [OR:B ion 9 57043] :Y1393	:g263395 acillus of 21): [RE:574 7] [PN:r	subtili from 15 405] [DI outative	BSUB(s) [I 98421 :dire Asp2	0009] OB:genpept-bct1] Lto 1807200.] ect] OB:genpept-bct1] OB:genpept-bct2] OB:genpept-bct2
ORF Name AI7503000990 24406563 cl 742	NTID 1790	<u>AAID</u>	NT LengthL	AA ength 16	ore	probability
Description	1750	3302				
NO-HIT						
ORF Name AI7503000990 24407313 f3 590	<u>NTID</u>	<u>AAID</u>	NT LengthL	AA ength	ore	probability
Description	1,21					
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> LengthL	<u>AA</u> ength	ore	probability
A17503000990_24407327_c3_1017	1792	5564	612	203 62	21 1	.2e-60
Description sp:[LN:PYRE_BACSU] [AC:P25972] [GN:P [DE:OROTATE PHOSPHORIBOSYLTRANSFERAS >pir:[LN:F69686] [AC:F69686:A30492] [CL:orotate phosphoribosyltransferas [OR:Bacillus subtilis] [EC:2.4.2.10] [AC:Z99112:AL009126] [PN:orotate pho biosynthesis] [OR:Bacillus subtilis] subtilis complete genome (section 9 name: pyrX] [SP:P25972] [LE:30299] [[LN:BACPYROP] [AC:M59757] [PN:OMP-PR [DB:genpept-bct2] [DE:Bacillus subti uracilpermease (pyrP), aspartate tra glutaminase of carbamyl phosphate sy (pyrAB), dihydroorotase dehydrogenas transferase (pyrE)genes, complete cd [DI:direct]	E, (OPRI [PN:ord e:orotat [DB:pin sphoribo [DB:ger of 21): RE:30949 PP trans lis ORFI nscarban nthetase e(pyrD),	c) (OP btate ce pho c1] >g bsyltr hpept- from e] [DI sferas LA (py mylase e (pyr	RTASE)] phosphor sphoribo p:[GI:e1 ansferas bct1] [E 1598421t :direct] e] [GN:p rR), put (pyrB), AA),carb decarbox	[SP:P25] ribosylt: syltrans 185148:(se] [GN:] C:2.4.2 to 180720 >gp:[G: yyrE] [OR tative me dihydro tamyl pho	972] ransf sfera g2633 pyrE] .10] 00.] I:g14 R:Bac embra corot cspha cyrF)	[DB:swissprot] Gerase,] [GN:pyrE] Ise homology] [929] [LN:BSUB0009] [FN:pyrimidine [DE:Bacillus [NT:alternate gene 3394] Fillus subtilis] Ine-bound ase (pyrC), te synthetase , and OMP-PRPP
ORF Name AI7503000990 24407760 cl 773	NTID	AAID	NT Langth	AA sco	ore	probability
	1793	5565	LengthL			
Description NO-HIT	1793	5565		o o		

ORF Name	NTID AAID LengthLength score probability				
AI7503000990_24407936_c2_896	1794 5566 1122 373 1380 4.3e-141				
KD PROTEIN IN FMT-SPOVM INTERGENIC REGIONS [AC:F69878] [PN:conserved hypothetical hypothetical protein HI0365] [OR:Bacilog: Sqp:[GI:e1185166:g2633947] [LN:BSUB00] [OR:Bacillus subtilis] [DB:genpept-bc:9 of 21): from 1598421to 1807200.] [NG:E50265] [RE:51356] [DI:direct] > gp [PN:YloN protein] [GN:yloN] [FN:unknown	ON] [OR:BACILLUS SUBTILIS] [DE:HYPOTHETICAL 41.6 GION] [SP:O34617] [DB:swissprot] >pir:[LN:F69878] al protein ylon] [GN:ylon] [CL:conserved llus subtilis] [DB:pir2] O9] [AC:Z99112:AL009126] [GN:ylon] [FN:unknown] t1] [DE:Bacillus subtilis complete genome (section T:similar to hypothetical proteins] [SP:O34617] :[GI:e323524:g2337804] [LN:BSY13937] [AC:Y13937] wn] [OR:Bacillus subtilis] [DB:genpept-bct1] m the spoVM region.] [SP:O34617] [LE:12037]				
	NTID AAID NT AA score probability 1795 5567 225 74				
ORF Name	NTID AAID NT AA score probability				
A17503000990_24415885_c3_960	1796 5568 1227 408 869 6.1e-87				
	carnosus condensing-enzyme-like protein (orf1) omplete cds.] [NT:orf2; unknown function; similar [DI:direct]				
ORF Name	NTID AAID LengthLength score probability				
A17503000990_24422077_c3_1013	1797 5569 1101 366 1196 1.4e-121				
Description sp:[LN:CARA_BACCL] [AC:P52557] [GN:PYRAA] [OR:BACILLUS CALDOLYTICUS] [EC:6.3.5.5] [DE:(EC 6.3.5.5) (CARBAMOYL-PHOSPHATE SYNTHETASE GLUTAMINE CHAIN)] [SP:P52557] [DB:swissprot] >pir:[LN:I40168] [AC:I40168:S34320] [PN:carbamoyl-phosphate synthase (glutamine-hydrolyzing),] [GN:pyrAa] [CL:carbamoyl-phosphate synthase (glutamine-hydrolyzing) small chain:carbamoyl-phosphate synthase (glutamine-hydrolyzing) small chain homology:trpG homology] [OR:Bacillus caldolyticus] [EC: 6.3.5.5] [DB:pir2] >gp:[GI:g312442] [LN:BCPYR] [AC:X73308] [PN:carbamoyl-phosphate synthase] [GN:PyrAa] [OR:Bacillus caldolyticus] [DB:genpept-bct1] [EC:6.3.5.5] [DE:B.caldolyticus pyrimidine biosynthesis genes.] [SP:P52557] [LE:2571] [RE:3665] [DI:direct]					
ORF Name	NTID AAID NT AA score probability				
Description pir: [LN:E69879] [AC:E69879] [PN:conse [CL:Mycoplasma genitalium hypothetical >gp:[GI:e1185175:g2633956] [LN:BSUB000 [OR:Bacillus subtilis] [DB:genpept-bct]	erved hypothetical protein yloV] [GN:yloV] l protein MG369] [OR:Bacillus subtilis] [DB:pir2] O9] [AC:Z99112:AL009126] [GN:yloV] [FN:unknown] t1] [DE:Bacillus subtilis complete genome (section T:similar to hypothetical proteins] [LE:57421]				

ORF Name AI7503000990_24475252_f2_371	NTID [1799	<u>AAID</u> 5571	NT Length	AA Length	score	probability		
Description NO-HIT			JLJ		J			
ORF Name	NTID	AAID	NT	<u>AA</u>	score	probability		
A17503000990_24484828_c2_832	1800	5572	1552	Length	200	8.2e-28		
Description	L	L	الـــــا ل		J L			
gp:[GI:g4981179] [LN:AE001739] [AC:AE001739:AE000512] [PN:conserved hypothetical protein] [GN:TM0656] [OR:Thermotoga maritima] [DB:genpept-bct2] [DE:Thermotoga maritima section 51 of 136 of the complete genome.] [NT:similar to SP:P38522 GB:U00096 PID:1742120] [LE:1379] [RE:1909] [DI:complement]								
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability		
A17503000990_24485950_c3_951	1801	5573	537	178	91	0.015		
Description gp:[GI:g5306139] [LN:AF160864] [AC:AF160864] [PN:NADH dehydrogenase subunit 2] [GN:nad2] [OR:Mitochondrion Tetrahymena pyriformis] [SR:Tetrahymena pyriformis] [DB:genpept] [EC:1.6.5.3] [DE:Tetrahymena pyriformis mitochondrial DNA, complete genome.] [LE:10108] [RE:10644] [DI:complement]								
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability		
A17503000990_24609637_c3_1001	1802	5574	798	265	916	6.4e-92		
Description gp:[GI:g4009492] [LN:AF068904] [AC:AF068904] [PN:YlmD] [GN:ylmD] [OR:Staphylococcus aureus] [DB:genpept-bct2] [DE:Staphylococcus aureus cell division protein FtsZ (ftsZ) gene,partial cds; YlmD (ylmD), YlmE (ylmE), YlmF (ylmF), YlmG (ylmG),and YlmH (ylmH) genes, complete cds; and cell division proteinDivIVA (divIVA) gene, partial cds.] [NT:similar to Bacillus subtilis YlmD] [LE:437] [RE:1228] [DI:direct]								
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability		
A17503000990_24610885_f2_352	1803	5575		140				
Description NO-HIT			1					
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability		
AI7503000990_24632827_c1_786	1804	5576	963	320	917	5.0e-92		
Description								
pir:[LN:A69626] [AC:A69626] [PN:meth [CL:methionyl-tRNA formyltransferase homology] [OR:Bacillus subtilis] [EC [LN:BSUB0009] [AC:Z99112:AL009126] [1 [OR:Bacillus subtilis] [DB:genpept-bounded and subtilis] [DB:genpept-bounded and subtilis] [DI:direct] >gr [LE:47978] [RE:48931] [DI:direct] >gr [PN:putative Fmt protein] [GN:yloL] [DB:genpept-bct1] [DE:Bacillus subtiling [RE:10703] [DI:direct]	: phosph :2.1.2.9 PN:methict1] [EC 21to 180 p:[GI:e3 [FN:unkr	noribos 9] [DB: ionyl-t C:2.1.2 07200.] 323503: nown] [ylglyc pir2] RNA fo .9] [D [NT:a g23378 OR:Bac	inamid >gp:[G rmyltr E:Baci lterna 02] [L	e form I:e118 ansfer llus s te gen N:BSY1 subtil	nyltransferase (5164:g2633945] (ase] [GN:fmt] (ubtilis complete (le name: yloL] (3937] [AC:Y13937] (is]		

AΑ NT ORF Name NTID AAID score probability LengthLength A17503000990 24642042 c2_853 1805 561 186 7.1e-22 Description pir: [LN:C69875] [AC:C69875] [PN:hypothetical protein ylbN] [GN:ylbN] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e334782:g2340010] [LN:BS16823KB] [AC:Z98682] [PN:YlbN protein] [GN:ylbN] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis genomic DNA 23.9kB fragment.] [LE:20393] [RE:20911] [DI:direct] >gp:[GI:e1185097:g2633878] [LN:BSUB0008] [AC:Z99111:AL009126] [GN:ylbN] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 8 of 21): from 1394791to 1603020.] [LE:179876] [RE:180394] [DI:direct] NTORF Name NTID AAID probability score LengthLength AI7503000990_24642817_c2_862 1806 5578 810 1.1e-121 Description sp:[LN:MURI STAHA] [AC:P52974] [GN:DGA] [OR:STAPHYLOCOCCUS HAEMOLYTICUS] [EC:5.1.1.3] [DE:GLUTAMATE RACEMASE,] [SP:P52974] [DB:swissprot] >gp:[GI:g520574] [LN:SHU12405] [AC:U12405] [PN:glutamate racemase] [GN:dga] [OR:Staphylococcus haemolyticus] [DB:genpept-bct1] [DE:Staphylococcus haemolyticus Y176 glutamate racemase (dga) gene, complete cds.] [LE:263] [RE:1063] [DI:direct] NT AANTID Length Length score ORF Name AAID probability A17503000990 24643836 c3 943 1807 5579 633 210 6.6e-19 Description pir:[LN:D69870] [AC:D69870:A36718] [PN:conserved hypothetical protein ykyA:hypothetical protein (aceA 5' region)] [GN:ykyA] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1185047:g2633828] [LN:BSUB0008] [AC:Z99111:AL009126] [GN:ykyA] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 8 of 21): from 1394791to 1603020.] [NT:alternate gene name: ykrC; similar to hypothetical] [LE:131900] [RE:132517] [DI:direct] >qp:[GI:q3282141] [LN:AF012285] [AC:AF012285:AF012284:U51911] [PN:unknown] [GN:ykrC] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct2] [DE:Bacillus subtilis mobA-nprE gene region.] [NT:similar to Orf5 encoded by GenBank Accession] [LE:32388] [RE:33005] [DI:direct] NT AΑ ORF Name NTID AAID score probability LengthLength A17503000990 24645025 c3 1012 1808 1284 427 3.1e-131 Description sp:[LN:PYRC BACCL] [AC:P46538] [GN:PYRC] [OR:BACILLUS CALDOLYTICUS] [EC:3.5.2.3] [DE:DIHYDROOROTASE, (DHOASE)] [SP:P46538] [DB:swissprot] >pir:[LN:I40167] [AC:I40167:S34319] [PN:dihydroorotase,] [GN:pyrC] [CL:Bacillus dihydroorotase:Bacillus dihydroorotase homology] [OR:Bacillus caldolyticus]

[EC:3.5.2.3] [DB:pir2] >gp:[GI:g312441] [LN:BCPYR] [AC:X73308] [PN:dihydroorotase] [GN:PyrC] [OR:Bacillus caldolyticus] [DB:genpept-bct1] [EC:3.5.2.3] [DE:B.caldolyticus

pyrimidine biosynthesis genes.] [SP:P46538] [LE:1285] [RE:2568] [DI:direct]

NTORF Name AAID NTID score probability LengthLength A17503000990 24648412 cl 798 1809 5581 133 .4e-27 Description sp:[LN:YLXM BACSU] [AC:P37104] [GN:YLXM] [OR:BACILLUS SUBTILIS] [DE:HYPOTHETICAL 13.2 KD PROTEIN IN FFH 5'REGION] [SP:P37104] [DB:swissprot] >pir:[LN:A47154] [AC:A47154:A69882] [PN:conserved hypothetical protein ylxM] [GN:ylxM] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1185188:q2633969] [LN:BSUB0009] [AC:Z99112:AL009126] [GN:ylxM] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 9 of 21): from 1598421to 1807200.] [NT:similar to hypothetical proteins] [LE:72807] [RE:73139] [DI:direct] >gp:[GI:d1023083:g2424968] [LN:D14356] [AC:D14356] [PN:ORF1] [GN:orf1] [OR:Bacillus subtilis] [SR:Bacillus subtilis (strain:168) DNA] [DB:genpept-bct1] [DE:Bacillus subtilis orf1, ffh, rpsP genes for ORF1, Ffh and 30Sribosomal protein S16, complete cds.] [LE:365] [RE:697] [DI:direct] NT ORF Name NTID AAID probability LengthLength AI7503000990 24652178 c2 847 1810 158 Description pir:[LN:F69930] [AC:F69930] [PN:conserved hypothetical protein yozB] [GN:yozB] [OR:Bacillus subtilis] [DB:pir2] >qp:[GI:e1185386;q2634307] [LN:BSUB0011] [AC:Z99114:AL009126] [GN:yozB] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 11 of 21): from 2000171to 2207900.] [NT:similar to hypothetical proteins] [LE:85155] [RE:85691] [DI:complement] NT AA ORF Name NTID AAID score probability LengthLength AI7503000990 24730340 c3 937 1811 5583 672 223 731 2.6e-72 Description sp:[LN:YKQB BACSU] [AC:P39760] [GN:YKQB] [OR:BACILLUS SUBTILIS] [DE:HYPOTHETICAL 24.3 KD PROTEIN IN KINC-ADEC INTERGENIC REGION (ORF4)] [SP:P39760] [DB:swissprot] >pir:[LN:A69862] [AC:A69862:PC6016] [PN:conserved hypothetical protein ykqB] [GN:ykqB [CL:conserved hypothetical protein MG323] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:d1007628:g520844] [LN:BACAMOKOOO] [AC:D37799] [PN:orf4] [OR:Bacillus subtilis] [SR:Bacillus subtilis (sub_species:Marburg, strain:168] [DB:genpept-bct1] [DE:Bacillus subtilis genes for ampS, mreBH, orf1, kinC, orf3, orf4 andorf5.] [LE:5175] [RE:5840] [DI:direct] >gp:[GI:e1185041:g2633822] [LN:BSUB0008] [AC:Z99111:AL009126] [GN:ykqB] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 8 of 21): from 1394791to 1603020.] [NT:alternate gene name: ylxV, yzaC: similar to] [SP:P39760] [LE:125146] [RE:125811] [DI:direct] >qp:[GI:q3282136] [LN:AF012285] [AC:AF012285:AF012284:U51911] [PN:unknown] [GN:ykqA] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct2] [DE:Bacillus subtilis mobA-nprE gene region.] [NT:similar to product of orf4 encoded by GenBank] [LE:25634] [RE:26299] [DI:direct] NT AΑ ORF Name NTID <u>score</u> probability LengthLength A17503000990_24801713_c3_946 1812 1314 437 1906 7.9e-197 Description sp:[LN:ODP2 STAAU] [AC:Q59821] [GN:PDHC] [OR:STAPHYLOCOCCUS AUREUS] [EC:2.3.1.12] [DE:COMPLEX, (E2)] [SP:Q59821] [DB:swissprot] >pir:[LN:S19722] [AC:S19722] [PN:dihydrolipoamide S-acetyltransferase, chain E2] [CL:dihydrolipoamide acetyltransferase: lipoyl/biotin-binding homology] [OR:Staphylococcus aureus] [EC:2.3.1.12] [DB:pir2] >gp:[GI:g581570] [LN:SAPDHDNA] [AC:X58434:S73625]

[PN:dihydrolipoamide S-acetyltransferase, chain E2] [CL:dihydrolipoamide acetyltransferase: lipoyl/biotin-binding homology] [OR:Staphylococcus aureus] [EC:2.3.1.12] [DB:pir2] >gp:[GI:g581570] [LN:SAPDHDNA] [AC:X58434:S73625] [PN:dihydrolipoamide acetyltransferase: subunit E2] [GN:pdhC] [OR:Staphylococcus aureus] [DB:genpept-bct1] [EC:2.3.1.12] [DE:S.aureus pdhB, pdhC and pdhD genes for pyruvate decarboxylase,dihydrolipoamide acetyltransferase and dihydrolipoamidedehydrogenase.] [SP:Q59821] [LE:557] [RE:1849] [DI:direct]

ORF Name	NTID	AAID	LengthLength score probability
A17503000990_24806587_c2_848	1813	5585	1056 351 399 3.9e-37
Description pir: [LN:H69873] [AC:H69873] [PN:con [OR:Bacillus subtilis] [DB:pir2] >gp [PN:YlbC protein] [GN:ylbC] [OR:Baci subtilis genomic DNA 23.9kB fragment >gp: [GI:e1185086:g2633867] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-b 8 of 21): from 1394791to 1603020.] [subtilis] [LE:170993] [RE:172033] [D	:[GI:e3 llus su .] [LE: 008] [A ct1] [D NT:simi	34771:9 btilis 11510] C:Z991 E:Baci lar to	g2339999] [LN:BS16823KB] [AC:Z98682] [DB:genpept-bct1] [DE:Bacillus [RE:12550] [DI:direct]
ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000990_24851577_c3_971 Description	1814	5586	564 187 405 9.0e-38
pir:[LN:E69874] [AC:E69874] [PN:con [OR:Bacillus subtilis] [DB:pir2] >gp [PN:YlbH protein] [GN:ylbH] [OR:Baci subtilis genomic DNA 23.9kB fragment >gp:[GI:e1185091:g2633872] [LN:BSUB0	:[GI:e3 llus su .] [LE: 008] [A ct1] [D	34776:g otilis] 14650] C:Z9911 E:Bacil	g2340004] [LN:BS16823KB] [AC:Z98682] [DB:genpept-bct1] [DE:Bacillus [RE:15144] [DI:direct] [FN:unknown] [Llus subtilis complete genome (section
ORF Name	NTID	AAID	NT AA LengthLength score probability
AI7503000990_25425202_c1_749	1815	5587	522 173 196 1.3e-15
Description pir:[LN:B69985] [AC:B69985] [PN:hypesubtilis] [DB:pir2] >gp:[GI:e1184109 [GN:yshB] [FN:unknown] [OR:Bacillus complete genome (section 15 of 21):[DI:complement] >gp:[GI:e1165328:g17 protein] [GN:yshB] [OR:Bacillus subt sequence 89009bp.] [NT:unknown funct	:g263533 subtilia from 27 70034] ilis] [1	25] [LM s] [DB: 95131tc [LM:BS2 DB:genr	N:BSUB0015] [AC:Z99118:AL009126] genpept-bct1] [DE:Bacillus subtilis 0 3013540.] [LE:129072] [RE:129605] Z75208] [AC:Z75208] [PN:hypothetical pept-bct1] [DE:B.subtilis genomic
ORF Name	NTID	AAID	NT AA score probability
Description pir:[LN:A27763] [AC:A27763:C69704] reductase:fumaric hydrogenase:succin			dehydrogenase, flavoprotein:fumarate
reductase flavoprotein:3-oxosteroid : flavoprotein homology] [OR:Bacillus :	l-dehyd:	rogenas	se homology:fumarate reductase
ORF Name	NTID	AAID	NT AA LengthLength score probability
AI7503000990_25509640_f1_185	1817	5589	240 75 0.0084
Description sp:[LN:YRUB_CLOPA] [AC:P23171] [OR:Cl [DB:swissprot] >pir:[LN:S29118] [AC:S] [OR:Clostridium pasteurianum] [DB:pir [OR:Clostridium pasteurianum] [SR:C.] [DB:genpept-bct1] [DE:C.pasteurianum] [gene,complete cds.] [NT:open reading	S29118] r2] >gp pasteur: open re	[PN:h [GI:gl Lanum (eading	sypothetical protein 2] .44907] [LN:CLORUB] [AC:M60116] strain ATCC 6013) DNA] frame A, B, C, and rubredoxin

ORF Name	NTID	AAID	Length Length	th score	probability
AI7503000990_25604677_c1_736	1818	5590	450 149	=	5.6e-29
Description pir:[LN:C69874] [AC:C69874] [PN:con [OR:Bacillus subtilis] [DB:pir2] >gp [PN:YlbF protein] [GN:ylbF] [OR:Baci subtilis genomic DNA 23.9kB fragment >gp:[GI:e1185089:g2633870] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-b 8 of 21): from 1394791to 1603020.] [[RE:173483] [DI:direct]	:[GI:e3 llus su .] [LE: 008] [A ct1] [D	34774: btilis 13551] C:Z991 E:Baci	g2340002] [] [DB:genpe [RE:14000] 11:AL009126 llus subtil	LN:BS168 pt-bct1] [DI:dir] [GN:yl is compl	[DE:Bacillus ect] [bF] [FN:unknown] ete genome (section
ORF Name	NTID	AAID	NT AA LengthLengt	score	probability
A17503000990_25635962_c3_1008	1819	5591	486 161	274	6.9e-24
Description gp:[GI:e244971:g1340128] [LN:SA1234] [DB:genpept-bct1] [DE:S.aureus orfs [DI:direct]		-			
ORF Name	NTID	AAID	NT AA LengthLengt	score	probability
A17503000990_25816552_c3_1004	1820	5592	777 258	984	4.0e-99
gene,partial cds; YlmD (ylmD), YlmE genes, complete cds; and cell divisi [NT:similar to Bacillus subtilis Ylm	on prot	einDiv	IVA (divIVA) gene,	partial cds.]
ORF Name	NTID	AAID	NT AA LengthLengt	h score	probability
A17503000990_25818811_c2_841	1821	5593	153 50	162	5.1e-12
Description gp:[GI:g1022725] [LN:SHU35635] [AC:U haemolyticus] [SR:Staphylococcus hae [DE:Staphylococcus haemolyticus IS12 [LE:394] [RE:1083] [DI:complement] > [OR:Staphylococcus aureus] [SR:Staphylococcus [DB:genpept-bct1] [DE:Staphylococcus geneand unknown ORF, complete cds.] [DI:complement]	molytic 72 ORF1 gp:[GI: ylococc aureus	us stra and O g29516 us auro methic	ain=Y176] [I RF2 genes, (2] [LN:STAM] eus (strain cillin-resi:	complete ECRA] [A COL) DN stance p	pt-bct1] cds.] [NT:ORF2] C:L14017] A] rotein (mecR)
ORF Name	NTID	AAID	NT AA LengthLengt	:h score	probability
A17503000990_25939030_t2_399 Description NO-HIT	1822	5594	[183] [60		
ORF Name					

	NTID	AAID <u>NT AA</u> score probability LengthLength
A17503000990_25976401_£3_533	1824	5596 150 49
Description	!	
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
AI7503000990_26354550_c1_740	1825	5597 192 63
Description		
NO-HIT		
ORF Name	NTID	AAID LengthLength score probability
A17503000990_26423305_c1_767	1826	5598 336 111 386 9.3e-36
<pre>aureus] [DB:genpept-bct2] [DE:Staphy gene,partial cds; YlmD (ylmD), YlmE</pre>	lococcus (ylmE), on prote	PN:YlmG] [GN:ylmG] [OR:Staphylococcus aureus cell division protein FtsZ (ftsZ) YlmF (ylmF), YlmG (ylmG),and YlmH (ylmH) einDivIVA (divIVA) gene, partial cds.]
ORF Name	NTID	AAID NT AA score probability
A17503000990 26597077 c2 875	1827	[5599 1398 465 1113 8.5e-113
Description		
	luster,	DE:Staphylococcus aureus strain ATCC yllB, yllC, yllD, pbpA, mraY, murD, div1B, [RE:8498] [DI:direct]
ORF Name	NTID	AAID LengthLength score probability
A17503000990_26598402_c3_1011	1828	5600 1314 437 1251 2.0e-127
Description sp:[LN:PYRP_BACCL] [AC:P41006] [GN:P (URACIL TRANSPORTER)] [SP:P41006] [D		
[PN:uracil transport protein:uracil [OR:Bacillus caldolyticus] [DB:pir2] [PN:uracil permease] [GN:pyrP] [OR:B	permease >gp:[G] acillus	e] [CL:uracil transport protein uraA] I:g431231] [LN:BCPYRQP] [AC:X76083]
[PN:uracil transport protein:uracil [OR:Bacillus caldolyticus] [DB:pir2] [PN:uracil permease] [GN:pyrP] [OR:B [DE:B.caldolyticus (DSM405) pyrR, py	permease >gp:[G] acillus	prot] >pir:[LN:S38893] [AC:S38893] e] [CL:uracil transport protein uraA] I:g431231] [LN:BCPYRQP] [AC:X76083] caldolyticus] [DB:genpept-bct1]
[PN:uracil transport protein:uracil [OR:Bacillus caldolyticus] [DB:pir2] [PN:uracil permease] [GN:pyrP] [OR:B [DE:B.caldolyticus (DSM405) pyrR, py [RE:2788] [DI:direct]	permease >gp:[G] acillus rP and p	prot] >pir:[LN:S38893] [AC:S38893] [E] [CL:uracil transport protein uraA] [I:g431231] [LN:BCPYRQP] [AC:X76083] [Caldolyticus] [DB:genpept-bct1] [PyrB (partial) genes.] [SP:P41006] [LE:1490]
[PN:uracil transport protein:uracil [OR:Bacillus caldolyticus] [DB:pir2] [PN:uracil permease] [GN:pyrP] [OR:B [DE:B.caldolyticus (DSM405) pyrR, py [RE:2788] [DI:direct] ORF Name A17503000990_26614167_c1_795 Description	permease >gp:[GI acillus rP and r	Prot] >pir:[LN:S38893] [AC:S38893] [E] [CL:uracil transport protein uraA] [I:g431231] [LN:BCPYRQP] [AC:X76083] [Caldolyticus] [DB:genpept-bct1] [PyrB (partial) genes.] [SP:P41006] [LE:1490] [AAID NT AA LengthLength score probability

ORF Name	NTID	AAID	NT AA score probability
AI7503000990_26828187_£1_147	1830	5602	
Description	<u> </u>		
<pre>complete genome (section 8 of 21): f [DI:complement] >gp:[GI:e324323:g222 [OR:Bacillus subtilis] [DB:genpept-b</pre>	:g263385 subtilis rom 1394 4771] [I	53] [Li s] [DB 4791to LN:BSZ E:Baci	N:BSUB0008] [AC:Z99111:AL009126] :genpept-bct1] [DE:Bacillus subtilis 1603020.] [LE:155412] [RE:155897] 97025] [AC:Z97025] [GN:ylaL]
ORF Name	NTID	AAID	NT AA LengthLength score probability
AI7503000990_2751260_f3_673	1831	5603	153 50
Description			
NO-HIT			
ORF Name	NTID	AAID	NT AA LengthLength score probability
AI7503000990_2766500_c3_938	1832	5604	153 50 113 2.8e-06
Description			
<pre>gp:[GI:g1022726] [LN:SHU35635] [AC:U haemolyticus] [SR:Staphylococcus hae [DE:Staphylococcus haemolyticus IS12 [LE:1101] [RE:1922] [DI:complement]</pre>	molyticu	ıs str	ain=Y176] [DB:genpept-bct1]
ORF Name	NTID	AAID	NT AA LengthLength score probability
AI7503000990_2789801_f2_436	1833	5605	366 121 163 1.2e-11
Description			-
pir:[LN:D70070] [AC:D70070] [PN:tra [CL:Bacillus subtilis probable trans [DB:pir2]	-		
ORF Name	NTID	AAID	NT AA score probability
A17503000990_2835285_f3_466	1834	5606	147 48
Description			
NO-HIT			
ORF Name	NTID	AAID	NT AA LengthLength
Ä17503000990_29319086_c1_739	1835	5607	519 172 467 2.4e-44
Description			
	re biosy g2340005 ilis] [C 208] [RE 008] [AC ct1] [DE	mthes: DB:genr C:15693 C:Z9911 C:Bacil	is protein kdtB] [OR:Bacillus :BS16823KB] [AC:Z98682] [PN:YlbI pept-bct1] [DE:Bacillus subtilis B] [DI:direct]

[LE:174691] [RE:175176] [DI:direct]

ORF Name AI7503000990_29320217_t2_271 Description NO-HIT	NTID 1836	AAID NT AA score probability 5608 207 68
<pre>carboxylase:biotin carboxylase homol subtilis] [DB:pir2] >gp:[GI:e1185076 [PN:pyruvate carboxylase] [GN:pycA]</pre>	ogy:lipo :g263385 [OR:Baci omplete	genome (section 8 of 21): from 1394791to
ORF Name A17503000990_29410908_c2_867 Description NO-HIT	NTID 1838	AAID NT AA score probability [5610 129 42
	OR:Kinet pept-inv t 7 (ND7	coplast Blastocrithidia culicis] [DE:Blastocrithidia culicis ATCC30268] [Ogene, partial cds, and cytochrome C
[SP:P94464] [DB:swissprot] >pir:[LN:homolog yloM] [GN:yloM] [CL:hypothe: [DB:pir2] >gp:[GI:e1185165:g2633946] [FN:unknown] [OR:Bacillus subtilis] genome (section 9 of 21): from 15984: similar to RNA-binding] [SP:P94464] >gp:[GI:e323504:g2337803] [LN:BSY139]	E69878] tical pr [LN:BSU [DB:genr 21to 180 [LE:4891 37] [AC:	JB0009] [AC:Z99112:AL009126] [GN:yloM] Dept-bct1] [DE:Bacillus subtilis complete J7200.] [NT:alternate gene name: sun; JRE:50261] [DI:direct] Y13937] [PN:putative Fmu protein] [GN:yloM] Dept-bct1] [DE:Bacillus subtilis genomic DNA

NT ORF Name NTID probability LengthLength A17503000990 30663955 c2 809 1841 5613 1152 383 3.9e-76 Description sp:[LN:PURK BACSU] [AC:P12045] [GN:PURK] [OR:BACILLUS SUBTILIS] [EC:4.1.1.21] [DE:(AIR CARBOXYLASE) (AIRC)] [SP:P12045] [DB:swissprot] >pir:[LN:DCBSPK] [AC:B29326:G69684] [PN:phosphoribosylaminoimidazole carboxylase, carbon dioxide-fixation chain:phosphoribosylaminoimidazole carboxylase chain II] [GN:purK] [CL:phosphoribosylaminoimidazole carboxylase carbon dioxide-fixation chain:phosphoribosylaminoimidazole carboxylase carbon dioxide-fixation chain homology] [OR:Bacillus subtilis] [EC:4.1.1.21] [DB:pir1] [MP:18 min] >qp:[GI:q143365] [LN:BACPURF] [AC:J02732:K00047] [OR:Bacillus subtilis] [SR:B.subtilis (strain DE1 (prototroph DER. or W168)) DNA, clone pPZ] [DB:genpept-bct1] [DE:B.subtilis pur operon encoding purine biosynthesis enzymes, 12genes.] [NT:phosphoribosyl aminoimidazole carboxylase II] [LE:920] [RE:2059] [DI:direct] >gp:[GI:e1182623:g2632957] [LN:BSUB0004] [AC:Z99107:AL009126] [PN:phosphoribosylaminoimidazole carboxylase II] [GN:purK] [FN:purine biosynthesis] [OR:Bacillus subtilis] [DB:genpept-bct1] [EC:4.1.1.21] [DE:Bacillus subtilis complete genome (section 4 of 21): from 600701 to813890.] [SP:P12045] [LE:97902] [RE:99041] [DI:direct] NT AA ORF Name NTID AAID score probability Length Length AI7503000990 30745680 f2 431 1842 5614 2007 668 Description sp:[LN:QOX1 BACSU] [AC:P34956] [GN:QOXB:IPA-38D] [OR:BACILLUS SUBTILIS] [EC:1.9.3.-] [DE:SUBUNIT QOXB) (OXIDASE AA(3) SUBUNIT 1)] [SP:P34956] [DB:swissprot] >pir:[LN:B38129] [AC:B38129:S39693:F69687] [PN:bo-type ubiquinol oxidase, chain I:cytochrome aa3 quinol oxidase chain I:quinol oxidase aa3-600] [GN:qoxB] [CL:cytochrome-c oxidase chain I:cytochrome-c oxidase chain I homology] [OR:Bacillus subtilis] [EC:1.10.3.-] [DB:pir2] >gp:[GI:g143397] [LN:BACQOXA] [AC:M86548] [PN:quinol oxidase] [GN:QOXB] [OR:Bacillus subtilis] [SR:Bacillus subtilis DNA] [DB:genpept-bct1] [DE:Bacillus subtilis AA3-600 quinol oxidase (QOXA, QOXB, QOXC, QOXD)genes, complete cds.] [LE:1846] [RE:3795] [DI:direct] >gp:[GI:g413962] [LN:BSGENR] [AC:X73124] [GN:ipa-38d qoxB] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:B.subtilis genomic region (325 to 333).] [SP:P34956] [LE:40298] [RE:42247] [DI:direct] >gp:[GI:e1186315:g2636351] [LN:BSUB0020] [AC:Z99123:AL009126] [PN:cytochrome aa3 quino1 oxidase (subunit I)] [GN:qoxB] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 20 of 21): from 3798401to 4010550.] [NT:alternate

 ORF Name
 NTID
 AAID
 NT AA Length Length
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 A17503000990_31272062_f1_183
 1843
 5615
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 Description
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gene name: ipa-38d] [SP:P34956] [LE:115715] [RE:117664] [DI:complement]

NO-HIT

438

ORF Name	NTID	AAID	NT LengthL	AA ength score	probability
A17503000990_32242890_f2_397	1844	5616	1701 [5	1986	2.6e-205
Description sp:[LN:YKQC_BACSU] [AC:Q45493] [GN:YKD PROTEIN IN ADEC-PDHA INTERGENIC F [AC:B69862] [PN:conserved hypothetical protein MG139] [OR:BacilLN:BSUB0008] [AC:Z99111:AL009126] [DB:genpept-bct1] [DE:Bacillus subtitation of the second	REGION] ical pro- illus sul [GN:ykqC ilis com nypothet: :g328213: nknown] illus sul	[SP:Q4 tein y btilis] [FN: plete e ical p [GN:yke btilis	5493] [D kqC] [GN] [DB:pi unknown] genome (roteins] :AF01228 qC] [FN: mobA-np	B:swissprot [:ykqC] [CL [:r2] >gp:[GI [OR:Bacill section 8 o [SP:Q45493 [SP:Q45493 [SP:Q45493 [OrE gene reg	<pre>] >pir:[LN:B69862] :conserved :e1185043:g2633824] us subtilis] f 21): from] [LE:127733] R:Bacillus ion.] [NT:similar</pre>
ORF Name	NTID	AAID	NT	AA ength score	probability
AI7503000990 32667138 £1 135	1845	5617		ength———	
Description		L			
NO-HIT					
ORF Name	NTID	AAID	NT	AA sgoro	probability
	MIID	AAID	LengthL		probability
A17503000990_32756_£3_601	1846	5618	1401 4	748	4.0e-74
Description					
pir:[LN:S62667] [AC:S62667] [PN:Nra					nce-associated
macrophage protein 1] [OR:Oryza sati	LVaj [SR	:, rice	el [ng:b	1r2] 	
ORF Name	NTID	AAID	NT Length Le	AA ength	probability
A17503000990_3314128_f2_238	1847	5619		7	
Description					
NO-HIT					
ORF Name	NTID	AAID	NT LengthLe	AA ength score	probability
A17503000990_33153_c2_834	1848	5620	900 2	99 454	5.8e-43
Description	·		JLJ L		···
gp:[GI:g4981938] [LN:AE001791] [AC:A	E001791	AE000!	512] [PN	:spermidine,	/putrescine ABC
transporter, permease] [GN:TM1377] [
[DE:Thermotoga maritima section 103 GB:L42023 SP:P45170 PID:1007357] [LE					[NT:similar to
			72] [DI.		
ORF Name	NTID	AAID	NT LengthLe	AA ength score	probability
A17503000990_33367325_c1_706	1849	5621	168 5	5	
Description				······································	
NO-HIT					

ORF Name	NTID	AAID NT AA score probability
A17503000990_33594187_c1_728	1850	5622 303 100 150 7.1e-10
[DB:genpept-bct2] [DE:Lactobacillus	rhamnos nal regu	PN:unknown] [OR:Lactobacillus rhamnosus] us 6-phospho-beta-glucosidase homolog lator homolog and surfacelocated protein [RE:>3603] [DI:complement]
ORF Name	NTID	AAID NT AA score probability
AI7503000990_33595178_£3_476	1851	5623 129 42
Description NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000990_33651636_c3_981	1852	5624 1776 591 1528 9.0e-157
[AC:Z99118:AL009126] [GN:yshC] [FN:u [DE:Bacillus subtilis complete genom [NT:similar to DNA polymerase beta] >gp:[GI:e1165329:g1770035] [LN:BSZ75	nknown] ne (sect [LE:127 5208] [A 3:genpep	C:Z75208] [PN:hypothetical protein] t-bctl] [DE:B.subtilis genomic sequence
[OR:Bacillus subtilis] [DB:pir2] >gp [PN:putative fibronectin-binding pro [DB:genpept-bct1] [DE:Bacillus subti [LE:6297] [RE:8015] [DI:complement] [AC:Z99112:AL009126] [GN:yloA] [FN:u [DE:Bacillus subtilis complete genom	o:[GI:e3] otein] [0 .lis pyr] >gp:[GI onknown] ne (sect:	E to yloA gene region.] [NT:protein A-like]
ORF Name A17503000990_34100626_t3_494 Description	NTID 1854	AAID NT AA score probability LengthLength 5626 159 52

NO-HIT

NT ORF Name NTID probability LengthLength A17503000990 34175686 cl 686 5627 513 1855 4.7e-48 Description sp:[LN:PUR6 BACSU] [AC:P12044] [GN:PURE] [OR:BACILLUS SUBTILIS] [EC:4.1.1.21] [DE:(EC 4.1.1.21) (AIR CARBOXYLASE) (AIRC)] [SP:P12044] [DB:swissprot] >pir:[LN:DEBSPE] [AC:A29326:D69684] [PN:phosphoribosylaminoimidazole carboxylase, catalytic chain:phosphoribosylaminoimidazole carboxylase chain I] [GN:purE] [CL:phosphoribosylaminoimidazole carboxylase catalytic chain:phosphoribosylaminoimidazole carboxylase catalytic chain homology] [OR:Bacillus subtilis] [EC:4.1.1.21] [DB:pir1] [MP:18 min] >qp:[GI:q143364] [LN:BACPURF] [AC:J02732:K00047] [OR:Bacillus subtilis] [SR:B.subtilis (strain DE1 (prototroph DER. or W168)) DNA, clone pPZ] [DB:genpept-bct1] [DE:B.subtilis pur operon encoding purine biosynthesis enzymes, 12genes.] [NT:phosphoribosyl aminoimidazole carboxylase I] [LE:439] [RE:927] [DI:direct] >gp:[GI:e1182622:g2632956] [LN:BSUB0004] [AC:Z99107:AL009126] [PN:phosphoribosylaminoimidazole carboxylase I] [GN:purE] [FN:purine biosynthesis] [OR:Bacillus subtilis] [DB:genpept-bct1] [EC:4.1.1.21] [DE:Bacillus subtilis complete genome (section 4 of 21): from 600701 to813890.] [SP:P12044] [LE:97421] [RE:97909] [DI:direct] NTAΑ ORF Name NTID AAID <u>sco</u>re probability <u>LengthLength</u> A17503000990_34407750_f2_368 1856 5628 162 53 119 1.8e-07 Description pir: [LN:G69872] [AC:G69872] [PN:hypothetical protein ylaI] [GN:ylaI] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1185069:g2633850] [LN:BSUB0008] [AC:Z99111:AL009126] [GN:ylaI] [FN:unknown] [OR:Bacillus subtilis] [DB:qenpept-bct1] [DE:Bacillus subtilis complete genome (section 8 of 21): from 1394791to 1603020.] [LE:153004] [RE:153213] [DI:complement] >gp:[GI:e324322:q2224768] [LN:BSZ97025] [AC:Z97025] [GN:ylaI] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis nprE, yla[A,B,C,D,E,F,G,H,I,J,K,L,M,N,O] and pycAgenes.] [LE:7263] [RE:7472] [DI:complement] NT AA ORF Name NTID AAID score probability LengthLength A17503000990 34642213 c2 877 1857 615 204 681 5.1e-67 Description gp:[GI:g4009494] [LN:AF068904] [AC:AF068904] [PN:YlmF] [GN:ylmF] [OR:Staphylococcus aureus] [DB:genpept-bct2] [DE:Staphylococcus aureus cell division protein FtsZ (ftsZ) gene, partial cds; YlmD (ylmD), YlmE (ylmE), YlmF (ylmF), YlmG (ylmG), and YlmH (ylmH) genes, complete cds; and cell division proteinDivIVA (divIVA) gene, partial cds.] [NT:similar to Bacillus subtilis YlmF] [LE:1926] [RE:2480] [DI:direct] NT AA ORF Name NTID AAID score probability LengthLength A17503000990 34644125_c2_876 1858 5630 1416 471 1792 9.5e-185 Description

sp:[LN:FTSA_STAAU] [AC:007325] [GN:FTSA] [OR:STAPHYLOCOCCUS AUREUS] [DE:CELL DIVISION PROTEIN FTSA] [SP:007325] [DB:swissprot] >gp:[GI:q2149897] [LN:SAU94706] [AC:U94706] [PN:cell division protein] [GN:ftsA] [OR:Staphylococcus aureus] [DB:genpept-bct2] [DE:Staphylococcus aureus strain ATCC 8325-4 cell wall/cell divisiongene cluster, yllB, yllC, yllD, pbpA, mraY, murD, divlB, ftsA andftsZ genes, complete cds.] [LE:8604] [RE:10019] [DI:direct]

ORF Name	NTID AAI	D NT A	- score	probability
A17503000990_34646937_c3_983	1859 563	1 336 111	512 4	.1e-49
Description gp:[GI:e1333201:g3776111] [LN:SATRXA [FN:thiol:disulfide interchange] [OR [DE:Staphylococcus aureus trxA and u [LE:2334] [RE:2648] [DI:direct]	:Staphyloco	ccus aureus]	[DB:genpe	ept-bct1]
ORF Name	NTID AAI	D NT A	- 60000	probability
A17503000990_34650452_c1_782	1860 563	2 225 74	87 0	.00078
Description gp:[GI:g488925] [LN:A13473] [AC:A134 [SR:malaria parasite P. falciparum] antigen, clone 41-14.] [LE:<1] [RE:>	[DB:genpept	-pat] [DE:P.		_
ORF Name	NTID AAI	D NT A	- score	probability
A17503000990_35285902_c2_859	1861 563	3 150 49	46 0	.042
[EC:1.14.99.1] [DE:SYNTHASE 2) (PGH [DB:swissprot] >gp:[GI:g2959708] [LN 2] [OR:Mustela vison] [SR:American m prostaglandin synthase 2 mRNA, compl [DI:direct]	:AF047841] ink] [DB:ge	[AC:AF047841 npept-mam] [] [PN:pros DE:Mustela	taglandin synthase vison
ORF Name	NTID AAI	D NT A	A gth score	probability
AI7503000990_353427_£2_302	1862 563			
Description NO-HIT				
ORF Name	NTID AAI	NT A		probability
A17503000990_35365635_c2_846	1863 563	5 156 51		
Description NO-HIT				
ORF Name	NTID AAI	NT AM Length Leng		probability
A17503000990_35947191_c2_835	1864 563	6 1155 384	149 6	.3e-07
Description				
gp:[GI:g1633572] [LN:KSU52064] [AC:U [SR:Kaposi's sarcoma-associated herp [DE:Kaposi's sarcoma-associated herp [NT:Herpesvirus saimiri ORF73 homolo [LN:KSU75698] [AC:U75698] [OR:Kaposi sarcoma-associated herpesvirus - Hum sarcoma-associated herpesvirus long complete cds.] [NT:ORF 73; extensive	esvirus - H es-like vir g] [LE:1] [s sarcoma- an herpesvi unique regi	uman herpesv us ORF73 hom RE:3489] [DI associated he rus 8] [DB:go on, 80putati	irus 8] [D olog gene, :direct] > erpesvirus enpept-vrl ve ORF's a	B:genpept-vrl] complete cds.] gp:[GI:g1718329]] [SR:Kaposi's] [DE:Kaposi's nd kaposin gene,

[RE:127297] [DI:complement]

ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	score	probability
A17503000990_35978392_c1_788	1865	5637				2.0e-95
Description pir:[LN:H69878] [AC:H69878] [PN:prokinase homology] [OR:Bacillus subtil [LN:BSUB0009] [AC:Z99112:AL009126] [DB:genpept-bct1] [DE:Bacillus subtil 1598421to 1807200.] [NT:similar to p >gp:[GI:e323506:g2337806] [LN:BSY139 [GN:yloP] [FN:unknown] [OR:Bacillus genomic DNA from the spoVM region.]	is] [DB GN:yloP] lis comp rotein } GR [AC: subtilis	:pir2] [FN:: plete cinase Y1393 [DB	>gp:[0 unknowr genome] [LE:5 7] [PN:	GI:e118 n] [OR: (secti G2121] :putati pt-bct1	Bacillon 9 of [RE:54].ve Pkr	g2633949] lus subtilis] of 21): from 4067] [DI:direct] n2 protein] :Bacillus subtilis
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000990_35980062_£2_357	1866	5638	162	53]	
Description NO-HIT			<u> </u>			
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
AI7503000990_36129451_c1_702	1867	5639	288	95	408	4.3e-38
[DB:swissprot] >pir:[LN:A42374] [AC:sphosphohistidine-containing protein [CL:phosphotransferase system phosphosystem phosphohistidine-containing protein [DB:pir1] >gp:[GI:g46908] [LN:SCPTSH] (HPr)] [GN:ptsH] [OR:Staphylococcus gene for histidine-containing protein	:phospho phistidi rotein h [AC:X6 carnosus	otrans: ne-cono nomolog 50766]	ferase ntainin gy] [OF [PN:Hi :genper	systemg prot R:Staph istidin ot-bct1	n HPr] ein:ph ylococ e-cont] [DE:	nosphotransferase ccus carnosus] caining protein :S.carnosus ptsH
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000990_36141893_c3_1039	1868		1248		1166	2.1e-118
Description gp:[GI:e1185186:g2633967] [LN:BSUB000] particle (docking protein)] [GN:ftsY] proteins] [OR:Bacillus subtilis] [DB genome (section 9 of 21): from 159842] [SP:P51835] [LE:71119] [RE:72108] [DE	[FN:in genpept 21to 180	volved: -bct1] 7200.]	d in se	ecretio Bacillu Alterna	n of e	extracellular cilis complete
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000990_36142817_c1_717	1869	5641	1149	382	925	7.1e-93
Description pir: [LN:A70180] [AC:A70180] [PN:sper protein (potA) homolog] [CL:ATP-bind: [SR:, Lyme disease spirochete] [DB:pi: [AC:AE001165:AE000783] [PN:spermidine: [OR:Borrelia burgdorferi] [SR:Lyme disease spirochete] [SR:Lyme disease spirochet	ing cass [r2] >gp e/putres [sease s e comple	ette h : [GI:g cine h :piroch te ger	nomolog g268856 ABC tra nete] [nome.]	y] [OR 2] [LN nsport DB:gen [NT:si	:Borre :AE001 er,] [pept-b milar	elia burgdorferi] .165] .GN:BB0642] .ct2] [DE:Borrelia to GB:M64519

ORF Name NTID AAID NT AA score probability
A17503000990_36147150_c1_757
Description
pir:[LN:C69986] [AC:C69986] [PN:conserved hypothetical protein ysnA] [GN:ysnA] [CL:Methanococcus jannaschii conserved hypothetical protein MJ0226] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1184085:g2635301] [LN:BSUB0015] [AC:Z99118:AL009126] [GN:ysnA] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 15 of 21): from 2795131to 3013540.] [NT:similar to hypothetical proteins] [LE:104515] [RE:105111] [DI:complement] >gp:[GI:e1165357:g1770060] [LN:BSZ75208] [AC:Z75208] [PN:hypothetical protein] [GN:ysnA] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:B.subtilis genomic sequence 89009bp.] [NT:homology to ORFo197 of Escherichia coli; unknown] [LE:64796] [RE:65392] [DI:direct]
ORF Name NTID AAID NT AA score probability
AI7503000990_36219187_c3_920 [1871] [5643] [2208] [735] [2062] [2.3e-213]
Description
sp:[LN:PURL_BACSU] [AC:P12042] [GN:PURL] [OR:BACILLUS SUBTILIS] [EC:6.3.5.3] [DE:SYNTHASE II)] [SP:P12042] [DB:swissprot] >pir:[LN:SYBS2G] [AC:G29326:C69685] [PN:phosphoribosylformylglycinamidine synthase, component II:formylglycinamide ribotide amidotransferase] [GN:purL] [CL:phosphoribosylformylglycinamidine synthase component II] [OR:Bacillus subtilis] [EC:6.3.5.3] [DB:pirl] [MP:18 min] >gp:[GI:g143369] [LN:BACPURF] [AC:J02732:K00047] [OR:Bacillus subtilis] [SR:B.subtilis (strain DE1 (prototroph DER. or W168)) DNA, clone pPZ] [DB:genpept-bct1] [DE:B.subtilis pur operon encoding purine biosynthesis enzymes, 12genes.] [NT:phosphoribosylformyl glycinamidine synthetase II] [LE:5060] [RE:7288] [DI:direct] >gp:[GI:e1182628:g2632962] [LN:BSUB0004] [AC:Z99107:AL009126] [PN:phosphoribosylformylglycinamidine synthetase I] [GN:purQ] [FN:purine biosynthesis] [OR:Bacillus subtilis] [DB:genpept-bct1] [EC:6.3.5.3] [DE:Bacillus subtilis complete genome (section 4 of 21): from 600701 to813890.] [SP:P12042] [LE:102042] [RE:104270] [DI:direct]
ORF Name NTID AAID NT AA score probability
A17503000990_36445263_£2_320 [1872] 5644 [147] 48
Description
NO-HIT
ORF Name NTID AAID NT AA score probability
17503000990_36520302_c2_870 1873 5645 189 62 250 2.4e-21
Description
ED. [IN.VIJE STANII] [AC.007319] [CN.VIJE] [OD.STADUVI OCOCCIIS AIDEIIS] [DE.UVDOTUETICAI

17.4 KD PROTEIN] [SP:007319] [DB:swissprot] >gp:[GI:g2149890] [LN:SAU94706] [AC:U94706] [PN:unknown] [GN:yllB] [OR:Staphylococcus aureus] [DB:genpept-bct2] [DE:Staphylococcus aureus strain ATCC 8325-4 cell wall/cell divisiongene cluster, yllB, yllC, yllD, pbpA, mraY, murD, div1B, ftsA andftsZ genes, complete cds.] [LE:537] [RE:971] [DI:direct]

ORF Name	NTID	AAID	Length I	<u>AA</u> Length	score	probability
A17503000990_36601687_c1_692	1874	5646	1260		1016	1.6e-102
Description		J	J L I	·		
sp:[LN:PUR2_BACSU] [AC:P12039] [GN:P [DE:RIBONUCLEOTIDE SYNTHETASE) (PHOS [DB:swissprot] >pir:[LN:AJBSAG] [AC: ligase,:phosphoribosylglycinamide sy [CL:phosphoribosylamineglycine lig [OR:Bacillus subtilis] [EC:6.3.4.13] [LN:AF011544] [AC:AF011544] [PN:phos [OR:Bacillus subtilis] [DB:genpept-b phosphoribosylaminoimidazole-carboxa cds,phosphoribosylglycinamide synthe (yecB), YecC (yecC), and YecD (yecD) synthase (yecE) gene, partialcds.] [[LE:263] [RE:1531] [DI:direct] >gp:[[OR:Bacillus subtilis] [SR:B.subtilide] [PPZ] [DB:genpept-bct1] [DE:B.subtilide] [DI:direct] >gp:[GI:e1182633:g263296] [PN:phosphoribosylglycinamide synthe [OR:Bacillus subtilis] [DB:genpept-bed] [OR:Bacillus subtilis] [DB:genpept-bed] [OR:Bacillus subtilis] [DB:genpept-bed] [OR:Bacillus subtilis] [DB:genpept-bed] [OR:Bacillus subtilis] [DB:genpept-bed] [OR:Bacillus subtilis] [DB:genpept-bed] [OR:Bacillus subtilis] [DB:genpept-bed]	PHORIBO B29183 nthetas ase: pl [DB:p: phoribo ct1] [I midefor tase (I genes, NT:ider GI:g143 s (stra s pur c mide sy 7] [LN: tase]	DSYLGLY :C69684 se] [GN hosphor ir1] [M DSYlgly DE:Baci rmyltra purD), ,comple ntified 3374] [ain DE1 Dperon yntheta :BSUB00 [GN:pur EC:6.3.	CINAMIDI [PN:] [:purD] ibosylar P:18 min cinamide llus sul nsferase YecA (ye te cds, by comp LN:BACPU (protot encoding se (PUR 04] [AC D] [FN:] 4.13] [I	E SYNT phosph mine n] >g e synt btilis e (pur ecA), and p plemen URF] [troph g puri -D; gt :Z9910 purine DE:Bac	glycing: [GI: hetase H-J) gutative tation AC:J02 DER. cone bic g] [LE 7:AL00 biosyillus	E)] [SP:P12039] Sylamineglycine ne ligase homology] (g2465561] e] [GN:purD] gene, partial (iveadenine deaminase (re glutamate n of E.coli TX267,] (2732:K00047] (or W168)) DNA, clone (osynthesis enzymes, (E:11971] [RE:13239] (o)9126] (onthesis] subtilis complete
ORF Name	NTID	AAID	NT	<u>AA</u>	score	probability
AI7503000990 3912890 f3 463	1875	5647	LengthI	Length 73		6.4e-21
Description	1073] [3047		/ 3	240	0.46-21
sp:[LN:RL28_BACST] [AC:P23374] [GN:R RIBOSOMAL PROTEIN L28] [SP:P23374] [[PN:ribosomal protein L28] [CL:Esche stearothermophilus] [DB:pir2]	DB:swis	ssprot]	>pir:[I	LN:A48	396]	[AC:A48396]
ORF Name	NTID	AAID	<u>NT</u> LengthI	<u>AA</u> Length	score	probability
A17503000990_3945257_£3_664	1876	5648	1176	391	738	4.6e-73
Description			<u> </u>			
pir:[LN:E69687] [AC:E69687:A38129:S3 II) qoxA:quinol oxidase aa3-600 chai chain II precursor: cytochrome-c oxi [DB:pir2] >gp:[GI:e1186316:g2636352] aa3 quinol oxidase (subunit II)] [GN [DE:Bacillus subtilis complete genom [NT:alternate gene name: ipa-37d] [L	n qoxA] dase cl [LN:BS ::qoxA] e (sect	[GN:qonain II GUB0020 GUR:Bao Gun 20	oxA] [0 homolog] [AC:Z9 cillus s of 21):	CL:bo- gy] [O 99123: subtil : from	type u R:Baci AL0091 is] [I 37984	ubiquinol oxidase .llus subtilis] .26] [PN:cytochrome DB:genpept-bct1] .01to 4010550.]
ORF Name	NTID	AAID	<u>NT</u> LengthL	<u>AA</u> ength	score	probability
A17503000990_3960881_f3_483	1877	5649	129	12		
Description NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> LengthL	AA ength	score	probability
AI7503000990_39808_£3_498	1878	5650	135	14		
Description						
NO-HIT						

ODE Name	NMTD.	3 3 TD	NT AA	2224	
ORF Name	NTID	AAID	<u>LengthLength</u>	score	probability
A17503000990_3992193_f3_496	1879	5651	177 58]	
Description					
NO-HIT		a transport and transport		Company of the Second S	
ORF Name	NTID	AAID	NT AA LengthLength	score	probability
AI7503000990_4067342_c3_977 Description	1880	5652	1086 361	1267 4	.1e-129
sp:[LN:SYFA_BACSU] [AC:P17921:P94539 [DE:-TRNA LIGASE ALPHA CHAIN) (PHERS >pir:[LN:YFBSA] [AC:H69675:I40459:S1 chain:phenylalanyl-tRNA synthetase a ligase alpha chain] [OR:Bacillus sub >gp:[GI:e1184113:g2635329] [LN:BSUBO synthetase (alpha subunit)] [GN:pheS [EC:6.1.1.20] [DE:Bacillus subtilis 3013540.] [SP:P17921] [LE:133410] [R >gp:[GI:e1165324:g1770030] [LN:BSZ75 beta subunit] [GN:pheS] [OR:Bacillus [DE:B.subtilis genomic sequence 8900 subunit] [SP:P17921] [LE:35463] [RE:)] [SP:F 1730] [1pha cha tilis] [015] [AC] [OR:Ba complete E:134444 208] [AC subtili 9bp.] [N	P17921 [PN:phain] [GEC:6. C:Z991 acillumon genor By July July July July July July July Jul	:P94539] [DB enylalanine GN:pheS] [CI 1.1.20] [DB:p 18:AL009126] s subtilis] me (section : :complement] 08] [PN:phenylalanyl-tRi	:swisspr -tRNA li L:phenyl pir1] [PN:phe [DB:genp 15 of 21 ylalany-	gase, alpha alaninetRNA mylalanyl-tRNA ept-bct1]): from 2795131to tRNA synthetase 6.1.1.20]
ORF Name	NTID	AAID	NT AA LengthLength	score	probability
AI7503000990_4103438_c3_970	1881	5653	255 84		.6e-06
Description		L	JL	J	
pir:[LN:D69874] [AC:D69874] [PN:con [OR:Bacillus subtilis] [DB:pir2] >gp [PN:YlbG protein] [GN:ylbG] [OR:Baci subtilis genomic DNA 23.9kB fragment >gp:[GI:e1185090:g2633871] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-b 8 of 21): from 1394791to 1603020.] [RE:173810] [DI:direct]	:[GI:e33 llus sub .] [LE:1 008] [AC ct1] [DE	84775: otilis .4055] C:Z991 :Baci	g2340003] [Li] [DB:genpept [RE:14327] 11:AL009126] llus subtilis	N:BS1682 -bct1] [DI:dire [GN:ylb s comple	3KB] [AC:Z98682] [DE:Bacillus ct] G] [FN:unknown] te genome (section
ORF Name	NTID	AAID	NT AA LengthLength	score	probability
A17503000990_4298177_c1_776	1882	5654	126 41	- 1	
Description			J	J	
NO-HIT					
ORF Name A17503000990_4300332_c1_700		<u>AAID</u> 5655	NT AA LengthLength	<u> </u>	probability
Description		L	/ L		
gp:[GI:g5306148] [LN:AF160864] [AC:A: Tetrahymena pyriformis] [SR:Tetrahymena pyriformis mitochondrial DNA, complete	ena pyri	formi	s] [DB:genpep	ot] [DE:	Tetrahymena

ATA initiation] [LE:15906] [RE:17003] [DI:complement]

ORF Name	NTID	AAID	<u>NT</u> LengthL	AA ength	ore	probability
A17503000990_4306562_c1_760	1884	5656	306 1	.01 46	9 1	5e-44
Description	····					
sp:[LN:YLLB_STAAU] [AC:007319] [GN:Y 17.4 KD PROTEIN] [SP:007319] [DB:swi [PN:unknown] [GN:yllB] [OR:Staphyloc aureus strain ATCC 8325-4 cell wall/mray, murD, div1B, ftsA andftsZ gene	ssprot] occus a cell di	>gp:[ureus] vision	GI:g2149 [DB:gen gene clu	890] [Li pept-bct ster, y]	N:SAU t2] llB,	J94706] [AC:U94706] [DE:Staphylococcus yllC, yllD, pbpA,
ORF Name	NTID	AAID	NT LengthLe	AA ength sco	ore	probability
A17503000990_4339053_c3_934	1885	5657	144 4	7		
Description NO-HIT			J			
ORF Name	NTID	AAID	NT Length Le	AA ength	ore	probability
A17503000990_4457788_f2_344	1886	5658	1716 5	71 28	5 5	.3e-21
Description						
pir:[LN:D71623] [AC:D71623] [PN:ery [GN:PFB0095c] [OR:Plasmodium falcip [AC:AE001371:AE001362] [PN:erythrocy [OR:Plasmodium falciparum] [SR:malar [DE:Plasmodium falciparum chromosome [NT:identified by sequence similarit [DI:complementJoin]	arum] [] te memb: ia para: 2, sec	DB:pir: rane p: site P tion 8	2] >gp:[0 rotein P: . falcipa of 73 o	GI:g3845 fEMP3] arum] [I f the co	5093] [GN:F OB:ge omple	[LN:AE001371] PFB0095c] enpept-inv2]
ORF Name	NTID	AAID	<u>NT</u> LengthLe	AA ength	ore	probability
A17503000990_4509661_c1_783	1887	5659	411 1	36 16	8 1	.2e-12
Description						
<pre>gp:[GI:e1453986:g4835313] [LN:SC5H1] [GN:SC5H1.10c] [OR:Streptomyces coel.coelicolor cosmid 5H1.] [NT:SC5H1.10c] [RE:8575] [DI:complement]</pre>	icolor]	[DB:ge	enpept-b	ct1] [DE	:Str	eptomyces
ORF Name	NTID	AAID	<u>NT</u> LengthLe	AA ength	ore	probability
AI7503000990_4532967_c3_982	1888	5660	2358 7	85 31	59 O	.0
Description		<u> </u>		—·		
gp:[GI:e1333200:g3776110] [LN:SATRXA] [FN:DNA mismatch repair protein] [OR [DE:Staphylococcus aureus trxA and umage: [RE:2161] [DI:direct]	:Staphy	lococci	ıs aureus	s] [DB:g	jenpe	pt-bct1]
ORF Name	NTID	AAID	<u>NT</u> LengthLe	AA ength	ore	probability
A17503000990_4696068_c1_741	1889	5661	186 6	1 23	7 5	.7e-20
Description						
sp:[LN:RL32_BACST] [AC:P07840] [GN:RI RIBOSOMAL PROTEIN L32 (RIBOSOMAL PROTEIN: >pir:[LN:R5BS37] [AC:S07236] [PN:ril BL37:ribosomal protein I] [CL:Escher: stearothermophilus] [DB:pir1]	TEIN I)	(BL37)] [SP:P(in L32:ri	07840] [ibosomal	DB:s	wissprot] tein

NT ORF Name NTID AAID score probability LengthLength A17503000990 4725000 cl 797 5662 3636 1211 Description pir:[LN:G69708] [AC:G69708:JC4819:PC4029] [PN:chromosome segregation SMC protein:minichromosome stabilizing protein SMC] [GN:smc] [CL:conserved hypothetical P115 protein] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1185185:g2633966] [LN:BSUB0009] [AC:Z99112:AL009126] [PN:chromosome segregation SMC protein homolg] [GN:smc] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 9 of 21): from 1598421to 1807200.] [NT:alternate gene name: ylqA] [LE:67539] [RE:71099] [DI:direct] NTAΑ LengthLength score ORF Name NTID AAID probability AI7503000990_4725415_c2_873 1891 5663 2337 3205 Description gp:[GI:d1023422:g2463561] [LN:AB007500] [AC:AB007500:D28879] [PN:penicillin-binding protein 1] [GN:pbpA] [OR:Staphylococcus aureus] [SR:Staphylococcus aureus (strain:NCTC8325) DNA] [DB:qenpept-bct1] [DE:Staphylococcus aureus genes for penicillin-binding protein 1, MraY, MurD, partial and complete cds.] [LE:594] [RE:2828] [DI:direct] >gp:[GI:g2149893] [LN:SAU94706] [AC:U94706] [PN:penicillin-binding protein 1] [GN:pbpA] [OR:Staphylococcus aureus] [DB:genpept-bct2] [DE:Staphylococcus aureus strain ATCC 8325-4 cell wall/cell divisiongene cluster, yllB, yllC, yllD, pbpA, mraY, murD, div1B, ftsA andftsZ genes, complete cds.] [LE:2318] [RE:4552] [DI:direct] NT AA ORF Name NTID AAID score probability LengthLength AI7503000990 4741010 c3 956 1892 5664 846 281 650 9.8e-64 Description sp:[LN:SUHB BACSU] [AC:Q45499] [GN:SUHB] [OR:BACILLUS SUBTILIS] [DE:EXTRAGENIC SUPPRESSOR PROTEIN SUHB HOMOLOG] [SP:Q45499] [DB:swissprot] >pir:[LN:E69864] [AC:E69864] [PN:myo-inositol-1(or 4)-monophosphatase homolog yktC] [GN:yktC] [CL:suppressor protein suhB] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1185057:g2633838] [LN:BSUB0008] [AC:Z99111:AL009126] [GN:yktC] [FN:unknown] [OR:Bacillus subtilis] [DB:qenpept-bct1] [DE:Bacillus subtilis complete genome (section 8 of 21): from 1394791to 1603020.] [NT:similar to myo-inositol-1(or 4)-monophosphatase] [SP:Q45499] [LE:142056] [RE:142853] [DI:direct] >gp:[GI:g3282150] [LN:AF012285] [AC:AF012285:AF012284:U51911] [PN:unknown] [GN:yktC] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct2] [DE:Bacillus subtilis mobA-nprE gene region.] [NT:similar to E. coli extragenic suppressor protein] [LE:42544] [RE:43341] [DI:direct] NT AΑ ORF Name NTID <u>score</u> probability LengthLength AI7503000990_4741703_c2_894 1893 5665 2418 805 5.3e-223 Description sp:[LN:PRIA BACSU] [AC:P94461:O34941] [GN:PRIA] [OR:BACILLUS SUBTILIS] [DE:PRIMOSOMAL PROTEIN N' (REPLICATION FACTOR Y)] [SP:P94461:034941] [DB:swissprot] >pir:[LN:A69682] [AC:A69682] [PN:primosomal replication factor Y priA] [GN:priA] [CL:unassigned DEAD/H box helicases:DEAD/H box helicase homology] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1185162:g2633943] [LN:BSUB0009] [AC:Z99112:AL009126] [PN:primosomal

replication factor Y (primosomal] [GN:priA] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 9 of 21): from 1598421to 1807200.] [NT:alternate gene name: yloJ] [SP:P94461] [LE:45047] [RE:47464] [DI:direct] >gp:[GI:e323502:g2337800] [LN:BSY13937] [AC:Y13937] [PN:putative PriA protein] [GN:yloJ] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis genomic DNA from the spoVM region.] [SP:P94461] [LE:6819] [RE:9236] [DI:direct]

ORF Name	NTID	AAID LengthLength score probability
A17503000990_4769512_t2_349	1894	[5666 [135]44
Description		
NO-HIT		
		NT AA
ORF Name	NTID	AAID LengthLength score probability
AI7503000990_4781517_c2_888	1895	5667 129 42
Description		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
A17503000990_4859425_£2_276	1896	5668 141 46
Description		
NO-HIT		
		NT AA
ORF Name	NTID	AAID LengthLength score probability
AI7503000990_4875055_f1_23	1897	5669 225 74 74 0.015
Description		
[GN:MJ0275.1] [OR:Methanococcus jannaschii section 24 of 150 of the	[SP:P812 1. janna naschii] complet	34] [DB:swissprot] >gp:[GI:g2826269] schii predicted coding region MJ0275.1] [DB:genpept-bct2] [DE:Methanococcus e genome.] [NT:Brute Force ORF; identified
by GeneMark; putative] [LE:9344] [RE	1:9/90]	[DI:Complement]
ORF Name	NTID	AAID NT AA core probability
ORF Name A17503000990_4875327_f3_587	NTID 1898	AAID NT AA score probability S670 135 44
		LengthLength score probability
A17503000990_4875327_f3_587 Description		LengthLength score probability [5670 135 44]
A17503000990_4875327_f3_587 Description NO-HIT	1898	LengthLength score probability 5670 135 44
A17503000990_4875327_f3_587 Description NO-HIT ORF Name	NTID	AAID LengthLength score probability AAID NT AA LengthLength score probability
Description NO-HIT ORF Name A17503000990_4875452_c3_1034 Description gp:[GI:g4982295] [LN:AE001811] [AC:A3-epimerase] [GN:TM1718] [OR:Thermot	NTID 1899 LE001811 loga mar:	AAID LengthLength score probability AAID NT AA LengthLength score probability 5671 645 214 521 4.6e-50 :AE000512] [PN:ribulose-phosphate itima] [DB:genpept-bct2] [DE:Thermotoga genome.] [NT:similar to GB:L77117 SP:Q58093
Description NO-HIT ORF Name A17503000990_4875452_c3_1034 Description gp:[GI:g4982295] [LN:AE001811] [AC:A 3-epimerase] [GN:TM1718] [OR:Thermot maritima section 123 of 136 of the comparison	NTID 1899 LE001811 loga mar:	AAID LengthLength score probability AAID NT AA LengthLength score probability 5671 645 214 521 4.6e-50 :AE000512] [PN:ribulose-phosphate itima] [DB:genpept-bct2] [DE:Thermotoga genome.] [NT:similar to GB:L77117 SP:Q58093
Description NO-HIT ORF Name A17503000990_4875452_c3_1034 Description gp:[GI:g4982295] [LN:AE001811] [AC:A 3-epimerase] [GN:TM1718] [OR:Thermot maritima section 123 of 136 of the c PID:1591395 percent] [LE:2607] [RE:3	NTID [1899] LE001811 loga mar complete [269] [D	AAID LengthLength score probability AAID NT AA LengthLength score probability 5671 645 214 521 4.6e-50 :AE000512] [PN:ribulose-phosphate itima] [DB:genpept-bct2] [DE:Thermotoga genome.] [NT:similar to GB:L77117 SP:Q58093 I:direct]
Description NO-HIT ORF Name A17503000990_4875452_c3_1034 Description gp:[GI:g4982295] [LN:AE001811] [AC:A 3-epimerase] [GN:TM1718] [OR:Thermot maritima section 123 of 136 of the c PID:1591395 percent] [LE:2607] [RE:3	NTID 1899 E001811 coga mar complete 269] [D	AAID LengthLength Secore probability AAID NT AA Score probability EngthLength Score probability EngthLength Score probability EngthLength Score probability AAID LengthLength Score probability EAE000512] [PN:ribulose-phosphate itima] [DB:genpept-bct2] [DE:Thermotoga genome.] [NT:similar to GB:L77117 SP:Q58093 I:direct] AAID NT AA LengthLength Score probability
Description NO-HIT ORF Name A17503000990_4875452_c3_1034 Description gp:[GI:g4982295] [LN:AE001811] [AC:A 3-epimerase] [GN:TM1718] [OR:Thermot maritima section 123 of 136 of the c PID:1591395 percent] [LE:2607] [RE:3 ORF Name A17503000990_4876542_f1_176 Description pir:[LN:D69862] [AC:D69862] [PN:for [CL:polypeptide deformylase] [OR:Bac >gp:[GI:e1185046:g2633827] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-b 8 of 21): from 1394791to 1603020.] [LE:130810] [RE:131364] [DI:compleme [AC:AF012285:AF012284:U51911] [PN:un subtilis] [DB:genpept-bct2] [DE:Baci	NTID 1899 LE001811 Loga mar: complete 269] [D: NTID 1900 mylmeth: cillus si constant Di NT:simi constant Sp known llus su	AAID LengthLength Score probability AAID LengthLength score probability EngthLength score probability EngthLength score probability EngthLength score probability AE000512] [PN:ribulose-phosphate itima] [DB:genpept-bct2] [DE:Thermotoga genome.] [NT:similar to GB:L77117 SP:Q58093 I:direct] AAID NT AA CORE Probability EngthLength score probability EngthLength scor

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
AI7503000990_4877203_c1_793	1901	5673		311	661	6.7e-65
Description pir: [LN:H69620] [AC:H69620] [PN:mal [GN:fabD] [CL:[acyl-carrier-protein S-malonyltransferase homology] [OR:B >gp:[GI:e1185181:g2633962] [LN:BSUB0 carrier protein transacylase] [GN:fa subtilis] [DB:genpept-bct1] [EC:2.3. (section 9 of 21): from 1598421to 18 [RE:65499] [DI:direct] >gp:[GI:e3235 FabD protein] [GN:ylpE] [FN:unknown] [DE:Bacillus subtilis genomic DNA fr [DI:direct]] S-mal acillus 009] [A bD] [FN 1.39] [07200.] 14:g233 [OR:Ba	onyltr subti C:Z991 :fatty DE:Bac [NT:a 7819] cillus	ansfera lis] [D 12:AL00 acid b illus s lternat [LN:BSY subtil	se: [a B:pir2 9126] iosynt ubtili e gene 13937] is] [D	cyl-ca [PN:ma hesis] s comp name: [AC:] B:genp	arrier-protein] alonyl CoA-acyl [OR:Bacillus blete genome ylpE] [LE:64546] (13937] [PN:putativ
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
AI7503000990_4899187_£3_569	1902	5674	1149	382	402	1.9e-37
Description pir: [LN:B69875] [AC:B69875] [PN:con [OR:Bacillus subtilis] [DB:pir2] >gp [PN:YlbM protein] [GN:ylbM] [OR:Baci subtilis genomic DNA 23.9kB fragment >gp: [GI:e1185096:g2633877] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-b 8 of 21): from 1394791to 1603020.] [[RE:179666] [DI:complement]	:[GI:e3 llus su .] [LE: 008] [A ct1] [D	34781: btilis 18936] C:Z991 E:Baci	g234000 [DB:g RE:20 11:AL00 llus su	9] [LN enpept 183] [1 9126] btilis	:BS168 -bct1] DI:com [GN:yl compl	B23KB] [AC:Z98682] [DE:Bacillus uplement] bM] [FN:unknown] ete genome (section
ORF Name	NTID	AAID	<u>NT</u> Length:	<u>AA</u> Length	score	probability
AI7503000990_4970462_c2_904	1903	5675		249	837	1.5e-83
Description sp:[LN:FABG_BACSU] [AC:P51831:O31733 [DE:ACYL CARRIER PROTEIN REDUCTASE)] [AC:A69621:PC4176] [PN:3-oxoacyl-[a carrier protein reductase fabG:srb p dehydrogenase:short-chain alcohol de [EC:1.1.1.100] [DB:pir2] >gp:[GI:e11 [PN:3-ketoacyl-acyl carrier protein [OR:Bacillus subtilis] [DB:genpept-b genome (section 9 of 21): from 15984 [SP:P51831] [LE:65492] [RE:66232] [D	[SP:P5 cyl-car rotein] hydroge: 85182:g: reducta: ct1] [E: 21to 18	1831:0 rier-p: [GN:fa nase ho 263396: se] [GI C:1.1.	31733] rotein] abG:srb omology 3] [LN: N:fabG]	[DB:sw: reduct] [CL] [OR:1 BSUB000 [FN:fa [DE:Bac	isspro tase,: :ribit Bacill 09] [A atty a cillus	ot] >pir:[LN:A69621 3-ketoacyl-acyl ol us subtilis] .C:Z99112:AL009126] .cid biosynthesis] subtilis complete
ORF Name A17503000990_5080260_f1_43 Description NO-HIT	NTID 1904	<u>AAID</u> 5676	<u>NT</u> Length]	AA Length	score	probability
ORF Name A17503000990_5096012_f3_481 Description	NTID 1905	<u>AAID</u> 5677	NT Lengthl	<u>AA</u> Length ⁵	score	probability

ORF Name	NTID	AAID	NT AA LengthLength score probability				
A17503000990_5117162_c2_826	1906	5678	1023 340 925 7.1e-93				
Description	·						
	tearothe] [DE:Ba II, comp	ermoph:	nilus] [SR:Bacillus stearothermophilus as stearothermophilus genes for bd-type				
ORF Name	NTID	AAID	NT AA LengthLength score probability				
A17503000990_5157963_£3_573	1907	5679	939 312 219 3.7e-22				
Description			ester phosphodiesterase homolog yhdW]				
[GN:yhdW] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1182963:g2633297] [LN:BSUB0006] [AC:Z99109:AL009126] [GN:yhdW] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 6 of 21): from 999501 to1209940.] [NT:similar to glycerophosphodiester phosphodiesterase] [LE:37668] [RE:38399] [DI:complement] >gp:[GI:e1191883:g2226218] [LN:BSY14082] [AC:Y14082] [PN:hypothetical protein] [GN:yhdW] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis chromosomal DNA, region 72 to 75 degrees: spoVRto sspB.] [NT:Similarity to glycerol diester phosphodiesterase] [LE:24316] [RE:25047] [DI:complement]							
ORF Name	NTID	AAID	NT AA score probability				
A17503000990_5162760_£3_514	1908	5680	129 42				
Description							
NO-HIT		_					
ORF Name	NTID	AAID	NT AA score probability				
A17503000990_5276712_c3_1016	1909	5681	708 235 659 1.1e-64				
Description							
sp:[LN:DCOP_LACLC] [AC:P50924] [GN:P [SR:,SUBSPCREMORIS:STREPTOCOCCUS CREI [SP:P50924] [DB:swissprot] >gp:[GI:e: decarboxylase] [GN:pyrF] [OR:Lactococ and pyrF genes.] [SP:P50924] [LE:5309	MORIS] [264705:g ccus lac	EC:4.1 107036 tis] [61] [LN:LLCPYRDB] [AC:X74207] [PN:OMP [DB:genpept-bct1] [DE:L.lactis pyrD				
ORF Name	NTID	AAID	NT AA score probability				
AI7503000990_595312_c1_733	1910	5682	147 48				
Description							
NO-HIT							
ORF Name AI7503000990 5974138 f1_158		<u>AAID</u> 5683	NT AA score probability				
Description	لــــــا		السيا لأسيا				

NO-HIT

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	proba	bility
A17503000990_5978453_c3_1029	1912	5684	492	163	261	1.6e-22	
Description							
sp:[LN:DEF_BACSU] [AC:P94462] [GN:DE [DE:DEFORMYLASE)] [SP:P94462] [DB:st [PN:polypeptide deformylase def] [GI subtilis] [DB:pir2] >gp:[GI:e290287 [PN:polypeptide deformylase] [GN:def [DE:B.subtilis priA, def, fmt, sun g >gp:[GI:e1185163:g2633944] [LN:BSUBG deformylase] [GN:def] [OR:Bacillus s [DE:Bacillus subtilis complete genor [NT:alternate gene name: yloK] [SP:I >gp:[GI:e323523:g2337801] [LN:BSY133] [FN:unknown] [OR:Bacillus subtilis] from the spoVM region.] [SP:P94462]	wissprot N:def] :g177249 f] [OR:E genes.] 0009] [A subtilis me (sect P94462] 937] [AC	[CL:pc [CL:pc [SP] [LN [SP:P9 [C:Z991]] [DB: [ion 9 [LE:47 [:Y1393]	E: [LN:F6 Plypepti I:BSPRIA IS SUBTI 14462] 12:ALOO genpept of 21): 491] [F 7] [PN:	G9613] de def ADFS] [lis] [[LE:176 9126] :-bct1] from RE:4797 putati	[AC:F6 ormyla AC:Y10 DB:gen 2] [RE [PN:pc [EC:3] [DI ve Defillus s	pept-bc :2244] olypepti :5.1.31 :lto 180 ::direct protei	t1] [DI:direct] de] 7200.]] n] [GN:yloK]
ORF Name	NTID	AAID	<u>NT</u> Length	<u> </u>	score		bility
A17503000990_6023593_c1_730 Description	1913	5685	288	95	296	3.2e-26	
pir: [LN:D69873] [AC:D69873] [PN:hypsubtilis] [DB:pir2] >gp:[GI:e1185074] [GN:ylaN] [FN:unknown] [OR:Bacillus complete genome (section 8 of 21): ff [DI:direct] >gp:[GI:e1191897:g222477] subtilis] [DB:genpept-bct1] [DE:Bacilyla[A,B,C,D,E,F,G,H,I,J,K,L,M,N,O] and subtilis]	1:g26338 subtili Erom 139 73] [LN: illus su	55] [L s] [DE 4791tc BSZ970 abtilis	N:BSUB0 :genpep 160302 25] [AC nprE,	0008] [bt-bct1 :0.] [L ::Z9702	AC:Z99] [DE: E:1570 5] [GN	111:AL0 Bacillu 27] [RE :ylaN]	09126] s subtilis :157308] [OR:Bacillus
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probal	bility
AY7503000990_6072125_c1_766	1914	5686		223	857	1.1e-85	
Description	,		J				
gp:[GI:g4009493] [LN:AF068904] [AC: aureus] [DB:genpept-bct2] [DE:Staphy gene,partial cds; YlmD (ylmD), YlmE genes, complete cds; and cell divisi [NT:similar to Bacillus subtilis Ylm	ylococcu (ylmE), ion prot	s aure YlmF einDiv	us cell (ylmF), IVA (di	divis YlmG vIVA)	ion pr (ylmG) gene,	otein Formand of the contract	tsZ (ftsZ) mH (ylmH)
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probab	oility
A17503000990_6115700_t3_647	1915	5687	1404	467	356	7.8e-64	
Description							
pir: [LN:H69858] [AC:H69858] [PN:catykoD] [GN:ykoD] [CL:ATP-binding casypp: [GI:e1181522:g2632042] [LN:BSAJ2subtilis] [DB:genpept-bct1] [DE:Baciand ykoR.] [NT:homologous to cobalt [RE:42578] [DI:complement] >gp:[GI:e[AC:Z99110:AL009126] [GN:ykoD] [FN:u[DE:Bacillus subtilis complete genoment]]	ssette h 2571] [A 1] llus su transpo 2] 183342 nknown] ne (sect	omolog C:AJ00 btilis rt ATP :g2633 [OR:B ion 7	y] [OR: 2571] [168 56 bindin 676] [L acillus of 21):	Bacillo PN:Ykol kb DN g proto N:BSUB subti from	us sub D] [GN A frag ein] [0007] lis] [119439	tilis] :ykoD] ment bet LE:41106 DB:genpe 1to 1411	[DB:pir2] [OR:Bacillus tween xlyA 6] ept-bct1]

[DI:complement]

ORF Name	NTID	AAID	NT AA score probability
A17503000990_6131693_c1_762	1916	5688	414 137 574 1.1e-55
Description		L	
[SR:Staphylococcus aureus (strain:NO aureus genes for penicillin-binding [NT:unnamed protein product] [LE:212 [LN:SAU94706] [AC:U94706] [PN:cell descriptions of the control of the contro	TC8325) protein [RE:61 livision lococcus lD, pbp	DNA] 1, Mr 13] [D prote s aure A, mra	DI:direct] >gp:[GI:g2149892] ein] [GN:yllD] [OR:Staphylococcus eus strain ATCC 8325-4 cell wall/cell
ORF Name A17503000990_6149077 c3 1033	NTID	<u>AAID</u>	NT AA LengthLength score probability [891 296 708 7.0e-70
Description	2327	3003	052 250 7.00 7.00 7.00
9 of 21): from 1598421to 1807200.] [RE:54978] [DI:direct] >gp:[GI:e3235	11714] 1009] [AG 10t1] [DI NT:simi] 125:g233 Bacillus	[OR:Ba C:Z991 E:Baci lar to 7807] s subt	acillus subtilis] [DB:pir2] 112:AL009126] [GN:yloQ] [FN:unknown] illus subtilis complete genome (section b hypothetical proteins] [LE:54082] [LN:BSY13937] [AC:Y13937] [PN:YloQ tilis] [DB:genpept-bct1] [DE:Bacillus
ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000990_632762_f3_671	1918	5690	<u> </u>
Description NO-HIT		1	
ORF Name	NTID	AAID	NT AA LengthLength
AI7503000990_642_c3_1005	1919	5691	2784 927 4356 0.0
Description sp:[LN:SYI_STAAU] [AC:P41972] [GN:IL [DE:(ILERS)] [SP:P41972] [DB:swisspr [PN:isoleucinetRNA ligase,:isoleuc [OR:Staphylococcus aureus] [EC:6.1.1]	ot] >pir yl-tRNA	c:[LN: synth	:S40178] [AC:S40178] netase] [CL:isoleucinetRNA ligase]
[AC:X74219] [PN:isoleucyl-tRNA synth	etase]	[GN:il	
[AC:X74219] [PN:isoleucyl-tRNA synth [DB:genpept-bct1] [DE:S.aureus gene	etase]	[GN:il	leS] [OR:Staphylococcus aureus]
[AC:X74219] [PN:isoleucyl-tRNA synth [DB:genpept-bct1] [DE:S.aureus gene [RE:2844] [DI:direct]	etase] for isol	[GN:ileleucyl	leS] [OR:Staphylococcus aureus] L-tRNA synthetase.] [SP:P41972] [LE:91] NT AA Garra probability
[AC:X74219] [PN:isoleucyl-tRNA synth [DB:genpept-bct1] [DE:S.aureus gene [RE:2844] [DI:direct] ORF Name	etase] for isol	[GN:il- Leucyl	leS] [OR:Staphylococcus aureus] L-tRNA synthetase.] [SP:P41972] [LE:91] NT AA LengthLength score probability

oxidase] [LE:363] [RE:1709] [DI:direct]

ORF Name	NTID	AAID	<u>NT</u> Length L	AA Length	core	probability
A17503000990_6720655_f3_530	1921	5693	147	48	68	0.045
Description						
<pre>gp:[GI:g4049913] [LN:AF063866] [AC:A [GN:MSV141] [OR:Melanoplus sanguinip sanguinipes entomopoxvirus, complete</pre>	es entor	vxoqom	irus] [I	DB:genp	ept-v	rl] [DE:Melanoplus
ORF Name	NTID	AAID	<u>NT</u> LengthL	AA Length	core	probability
A17503000990_6834687_c2_831	1922	5694	309	102	209	5.3e-17
Description						
<pre>pir:[LN:C69864] [AC:C69864] [PN:hyp subtilis] [DB:pir2] >gp:[GI:e1185054 [GN:yktA] [FN:unknown] [OR:Bacillus complete genome (section 8 of 21): f [DI:direct] >gp:[GI:g3282148] [LN:AF [GN:yktA] [FN:unknown] [OR:Bacillus mobA-nprE gene region.] [LE:41039] [</pre>	:g263383 subtilis rom 1394 012285] subtilis	35] [Li s] [DB 4791to [AC:A: s] [DB	N:BSUB00 :genpept 1603020 F012285: :genpept	008] [A :-bct1] D.] [LE :AF0122 :-bct2]	C:Z99 [DE: :1405 84:U5	111:AL009126] Bacillus subtilis 51] [RE:140817] 1911] [PN:unknown]
ORF Name	NTID	AAID	NT Length L	AA ength ^S	core	probability
A17503000990_7225000_c3_1038	1923	5695	1026	341	898	5.2e-90
Description			J			
<pre>>gp:[GI:e1185180:g2633961] [LN:BSUB0 fatty acid/phospholipid synthesis] [[DE:Bacillus subtilis complete genom [NT:alternate gene name: ylpD] [LE:6 >gp:[GI:e323529:g2337818] [LN:BSY139 [GN:ylpD] [FN:unknown] [OR:Bacillus genomic DNA from the spoVM region.]</pre>	OR:Bacil e (secti 3526] [F 37] [AC: subtilis	llus s ion 9 d RE:645 :Y1393 3] [DB	ubtilis] of 21): 27] [DI: 7] [PN:p :genpept	[DB:g from 1 direct outativ c-bct1]	enpep 59842] e Pls: [DE:]	t-bct1] 1to 1807200.] X protein] Bacillus subtilis
ORF Name	NTID	AAID	NT LengthL	AA ength	core	probability
AI7503000990_7320465_c1_719	1924	5696	831 2	276	525	1.7e-50
Description						
pir:[LN:G70179] [AC:G70179] [PN:sperprotein (potC) homolog] [CL:spermiding potI] [OR:Borrelia burgdorferi] [SR: >gp:[GI:g2688564] [LN:AE001165] [AC:transporter, permease] [GN:BB0640] [Ospirochete] [DB:genpept-bct2] [DE:Bocomplete genome.] [NT:similar to GB:I [RE:11098] [DI:complement]	ne/putre , Lyme d AE001165 OR:Borre rrelia b	escine disease 5:AE000 elia bu burgdon	transpo spiroc 1783] [P urgdorfe rferi (s	ort syschete] PN:sperieri] [Sisection	tem pe [DB:p: midine R:Lyme 51 of	ermease protein ir2] e/putrescine ABC e disease f 70) of the
ORF Name	NTID	AAID	<u>NT</u> LengthL	AA ength s	core	probability
A17503000990_867176_c3_924	1925	5697			090 2	2.3e-110
Description						
pir: [LN:C70468] [AC:C70468] [PN:phosformyltransferase] [GN:purH] [CL:put [DB:pir2] >gp:[GI:g2984204] [LN:AE006] [PN:phosphoribosylaminoimidazolecarbot [DB:genpept-bct2] [DE:Aquifex aeolicut [LE:7599] [RE:9119] [DI:complement]	rH bifur 0765] [A oxamide]	nctiona AC:AE00 [GN:p	al enzym 00765:AE purH] [O	ne] [OR :000657] :R:Aqui	:Aquii] fex a	fex aeolicus] eolicus]

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability	
A17503000990_892141_£2_303	1926	5698	396	131	281	1.2e-24	
Description gp:[GI:e1487496:g5042304] [LN:MMU242 (P4(21)n)] [GN:P4(21)n] [OR:Mus musculus partial mRNA for hypotheticamyh_yeast glucoamylase S1/S2] [LE:<[LN:AB028868] [AC:AB028868] [GN:P4(2:DB:genpept] [DE:Mus musculus P4(21)); with AMYH_YEAST GLUCOAMYLASE] [LE:<1	ulus] [S al prote 1] [RE:] 1)n] [OI n mRNA,	SR:hou ein (P 1203] R:Mus (partia	se mous 4(21)n [DI:dir musculu al cds.	e] [DE gene). ect] > s] [SF] [NT:	B:genpe] [NT: gp:[G] L:Mus m	ept-rod] [DE:Mus similar with ::d1042681:g5103287] nusculus mRNA]	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability	
A17503000990_892312_c3_980	1927	5699	156	51			
Description NO-HIT							
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability	
AI7503000990_968785_f2_361	1928	5700	939	312	1274	7.4e-130	
Description gp:[GI:g3320606] [LN:AF072726] [AC:AN [OR:Staphylococcus aureus] [DB:genpents genthase (ctaA) gene,complete cds.]	pt-bct2]	[DE:	Staphyl	ococcu	s aure	us putative heme A	
ORF Name	NTID	AAID	<u>NT</u> Lengthl	<u>AA</u> Length	score	probability	
AI7503000990_968811_c1_753	1929	5701	651	216		2.5e-58	
Description sp:[LN:DHSC_BACSU] [AC:P08064] [GN:SDHC] [OR:BACILLUS SUBTILIS] [DE:SUCCINATE DEHYDROGENASE CYTOCHROME B-558 SUBUNIT] [SP:P08064] [DB:swissprot] >pir:[LN:DEBSSC] [AC:A29843:I39972:E69704] [PN:succinate dehydrogenase, cytochrome b558:fumarate reductase C protein] [GN:sdhC:sdhA] [CL:succinate dehydrogenase cytochrome b558] [OR:Bacillus subtilis] [EC:1.3.99.1] [DB:pir1] [MP:70] >gp:[GI:g143525] [LN:BACSDHAB] [AC:M13470:M15107] [OR:Bacillus subtilis] [SR:B.subtilis DNA, clone pKIM4] [DB:genpept-bct1] [DE:B.subtilis succinate dehydrogenase complex encoding cytochromeb-558 subunit, complete cds, and flavoprotein subunit, 5' end.] [NT:succinate dehydrogenase cytochrome b-558 subunit] [LE:265] [RE:873] [DI:direct] >gp:[GI:e1184094:g2635310] [LN:BSUB0015] [AC:Z99118:AL009126] [PN:succinate dehydrogenase (cytochrome b558] [GN:sdhC] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 15 of 21): from 2795131to 3013540.] [SP:P08064] [LE:112102] [RE:112710] [DI:complement] >gp:[GI:e1165347:g1770051] [LN:BSZ75208] [AC:Z75208] [PN:cytochrome b558 subunit of succinate] [GN:sdhC] [OR:Bacillus subtilis] [DB:genpept-bct1] [EC:1.3.99.1] [DE:B.subtilis genomic sequence 89009bp.] [SP:P08064] [LE:57197] [RE:57805] [DI:direct]							
ORF Name	NTID	AAID	<u>NT</u> Length I	<u>AA</u> Length	score	probability	
A17503000990_969052_f2_434 Description NO-HIT	1930	5702	129	42			
ORF Name A17503000990_9881927_f1_75	NTID	<u>AAID</u> 5703	NT LengthI	AA Length	score	probability	
Description NO-HIT							

ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	score	probability
AI7503000990_990702_c1_765	1932	5704	1185	394	1809	1.5e-186
Description	LJ	<u> </u>	J L		. L	
sp:[LN:FTSZ_STAAU] [AC:P45498] [GN:FPROTEIN FTSZ] [SP:P45498] [DB:swisspdivision protein ftsZ] [CL:cell divi[DB:pirl] >gp:[GI:g458428] [LN:SAU06] [OR:Staphylococcus aureus] [DB:genpegene, complete cds.] [LE:27] [RE:119] [AC:U94706] [PN:cell division proteith [DB:genpept-bct2] [DE:Staphylococcus divisiongene cluster, yllB, yllC, ylcomplete cds.] [LE:10052] [RE:11224]	rot] >p sion pro 462] [A pt-bct1 9] [DI: 0 [GN: aureus 1D, pbp	ir:[LN otein : C:U0640] [DE:: direct] ftsZ] strain A, mra	:S58814 EtsZ] [PN Staphyl >gp:[[OR:Sta n ATCC	[AC: OR:Sta I:FtsZ] ococcu GI:g21 phyloc 8325-4	S58814 phylod [GN:fis aure 49898] coccus	PN:cell coccus aureus] ftsZ] eus SA4 FtsZ (ftsZ) [LN:SAU94706] aureus] wall/cell
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000990_9923437_f2_345	1933	5705		46		
Description		L		L		
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000990_9925910_c2_813	1934	5706	597	198	430	2.0e-40
Description						
sp:[LN:PUR3_BACSU] [AC:P12040] [GN:P [DE:TRANSFORMYLASE) (5'-PHOSPHORIBOS [DB:swissprot] >pir:[LN:XYBSGF] [AC: formyltransferase,] [GN:purN] [CL:p phosphoribosylglycinamide formyltran [EC:2.1.2.2] [DB:pir1] [MP:18 min] [OR:Bacillus subtilis] [SR:B.subtili pPZ] [DB:genpept-bct1] [DE:B.subtili 12genes.] [NT:phosphoribosyl glycina [DI:direct] >gp:[GI:e1182631:g263296 [PN:phosphoribosylglycinamide formyl [OR:Bacillus subtilis] [DB:genpept-b genome (section 4 of 21): from 60070 [DI:direct]	YLGLYCI) I29326:1 hosphor: sferase >gp:[GI s (stra: s pur op mide for 5] [LN:1 transfer ct1] [E0	NAMIDE B69685 ibosylo homolo :g14337 in DE1 peron e rmyltre BSUB000 rase] C:2.1.2	TRANSF [PN: glycina pgy] [O 72] [LN (proto encodin ansfera 04] [AC [GN:pur	ORMYLA phosph mide f R:Baci :BACPU troph g puri se] [L :Z9910 N] [FN E:Baci	oribos ormylt llus s RF] [A DER. c ne bic E:9833 7:AL00 :purin llus s	[SP:P12040] sylglycinamide cransferase: subtilis] AC:J02732:K00047] or W168)) DNA, clone osynthesis enzymes, B] [RE:10420] 09126] ne biosynthesis] subtilis complete
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	<u>score</u>	probability
AI7503000990_9933463_c3_933	1935	5707	1188	395	897	6.6e-90
Description sp:[LN:YWBD_BACSU] [AC:P39587] [GN:Y [DE:HYPOTHETICAL 44.4 KD PROTEIN IN [DB:swissprot] >pir:[LN:S39674] [AC: protein ipa-19d] [GN:ywbD] [CL:prob subtilis] [DB:pir2] >gp:[GI:g413943] subtilis] [DB:genpept-bct1] [DE:B.su [LE:20551] [RE:21741] [DI:direct] >g [AC:Z99123:AL009126] [GN:ywbD] [FN:u	EPR-GALI S39674:I able met [LN:BSC btilis c p:[GI:el	K INTER 370051 hyltra GENR] genomic L186335	RGENIC] [PN: nsfera [AC:X73 c regio 5:g2636	REGION ywbD p se b09 124] [n (325 371] [] [SP: rotein 67] [C GN:ipa to 33 LN:BSU	P39587] 1:hypothetical 1:R:Bacillus 1-19d] [OR:Bacillus 13).] [SP:P39587] 1B0020]

[DE:Bacillus subtilis complete genome (section 20 of 21): from 3798401to 4010550.] [NT:alternate gene name: ipa-19d; similar to] [SP:P39587] [LE:136218] [RE:137408]

[DI:complement]

ORF Name	NTID	AAID	NT AA score probability
A17503000990_9954640_£3_487	1936	5708	
Description			
<pre>gp:[GI:g2196686] [LN:EFU25090] [AC:U [GN:pyrAb] [OR:Enterococcus faecalis plasmid pKV48 pyrimidine biosynthesi [RE:>253] [DI:direct]</pre>] [DB:ge	enpept	-bct1] [DE:Enterococcus faecalis
ORF_Name	NTID	AAID	NT AA score probability
A17503000991_10928_f1_70	1937	5709	1242 413 238 1.4e-17
Description			
pir:[LN:T03492] [AC:T03492] [PN:hype [DB:pir2] [MP:1] >gp:[GI:g3128293] protein] [OR:Rhodobacter capsulatus] strain SB1003, partial genome.] [LE:	[LN:AF01 [DB:ger	10496] npept-l	[AC:AF010496] [PN:hypothetical bct2] [DE:Rhodobacter capsulatus
ORF Name	NTID	AAID	NT AA score probability
AI7503000991_10948587_c1_430	1938	5710	
Description			
<pre>gp:[GI:g4160468] [LN:AF109909] [AC:A [OR:Bacillus megaterium] [DB:genpept gene cluster, completesequence.] [NT [LE:277] [RE:1092] [DI:direct]</pre>	-bct2]	[DE:Ba	cillus megaterium polyhydroxyalkanoate
ORF Name	NTID	AAID	NT AA Score probability
A17503000991_11210760_f2_261	1939	5711	 162 53
Description		<u></u>	
NO-HIT			
ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000991_11220301_c2_523	1940	5712	126 41 108 3.4e-06
Description			
pir:[LN:QQSA8T] [AC:A04493] [PN:hype [DB:pir1]	othetica	al pro	tein B-189] [OR:Staphylococcus aureus]
ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000991_1180328_c3_606	1941	5713	816 271 372 2.8e-34
Description			
pir:[LN:H70003] [AC:H70003] [PN:prolaminopeptidase V:Pro-X aminopeptidase [GN:ytxM] [OR:Bacillus subtilis] [EG [LN:BSUB0016] [AC:Z99119:AL009126] [OB:genpept-bct1] [DE:Bacillus subtilized [DB:genpept-bct1] [DE:Bacillus subtilized [RE:151885] [DI:complement] >gp:[GI:GN:ytxM] [OR:Partillus subtilized [DB:genpept-bct1] [DI:complement] >gp:[GI:genpept-bct1] [DI:complement]]	e:prolin C:3.4.11 GN:ytxM] lis comp ne name: g2293147	e amin 5] [I 5] [I 	nopeptidase:proline iminopeptidase] DB:pir2] >gp:[GI:e1185954:g2635565] unknown] [OR:Bacillus subtilis] genome (section 16 of 21): from ; similar to prolyl] [LE:151061] :AF008220] [AC:AF008220] [PN:YtxM]
[GN:ytxM] [OR:Bacillus subtilis] [DB genomic region.] [NT:similarity with			

[RE:29366] [DI:direct]

ORF Name	NTID	AAID	NT AA LengthLengt	h score	probability
AI7503000991_12239817_c2_520	1942	5714	168 55	<u> </u>	
Description		L			
NO-HIT					
ORF Name	NTID	AAID	NT AA LengthLengt	h score	probability
A17503000991_12681510_f3_365	1943	5715	825 274	702 3	.0e-69
Description pir: [LN:F69841] [AC:F69841] [PN:conformation of the pir: [LN:F69841] [AC:F69841] [PN:conformation of the pir: [CN:Bacillus subtilis] [DB:pir: 2] > gp [GN:putative orf] [FN:unknown] [OR:Banger of the pir: 2] > gp [GN:putative orf] [DE:Bacillus subtilition of the pir: 2] [CN:BSUB0006] [AC:Z99109:AL009126] [GN:genpept-bct1] [DE:Bacillus subtilition of the pir: 2] [CN:Complement] > gp:[GI:e1173548:g214] [CN:Bacillus subtilis] [DB:genpept-bct] [CN:putative ORF] [LE:47689] [RE:4856] [RE:	:[GI:e26 acillus compleme GN:yitU] lis comp cal prot 45415]	subtient] > [FN: Tolete (cins] [LN:BS]	g1620926] [L lis] [DB:gen gp:[GI:e1183 unknown] [OR genome (sect [LE:190463] Y09476] [AC: btilis 54kb	N:BS168N pept-bct 116:g263 :Bacillu ion 6 of [RE:191 Y09476]	PRB] [AC:Z79580] 1] [DE:B.subtilis 3450] s subtilis] 21): from 999501 275] [PN:YitU]
ORF Name A17503000991_12750290_c1_400 Description pir:[LN:B69669] [AC:B69669:S15233:D38 protein oppD:sporulation initiation p membrane protein malK:ATP-binding cas >gp:[GI:e1183166:g2633500] [LN:BSUB08 transporter (ATP-binding] [GN:oppD] [OR:Bacillus subtilis] [DB:genpept-b8 7 of 21): from 1194391to 1411140.] [RE:29613] [DI:direct]	orotein ssette h 007] [AC [FN:requ ct1] [DE	spo0Kl nomolog :Z991: nired : :Baci	D] [GN:spo0K gy] [OR:Baci 10:AL009126] for initiati llus subtili	ransport D:oppD] llus sub [PN:oligon of spo	[CL:inner tilis] [DB:pir2] gopeptide ABC orulation,] te genome (section
ORF Name AI7503000991_13089052_f3_290	NTID 1945	<u>AAID</u> 5717	NT AA LengthLengtl 240 79		probability
Description pir:[LN:A60998] [AC:A60998] [PN:rep] [DB:pir2] >gp:[GI:g295834] [LN:SAPOX2 [DB:genpept-bct1] [DE:S.aureus plasm: [DI:direct]	2000] [A	C:X55	798] [OR:Sta	phylococo	cus aureus]
ORF Name A17503000991_13829403_f1_46 Description	NTID 1946	<u>AAID</u> 5718	NT AA LengthLengtl	score	probability

NO-HIT

ORF Name	NTID	<u>AAID</u>	LengthLength score probability		
A17503000991_13848387_t2_141	1947	5719	79 131 19.8e-09		
Description					
<pre>pir:[LN:S75993] [AC:S75993] [PN:hyp 6803, , PCC 6803] [SR:PCC 6803,] [D [AC:D64006:AB001339] [PN:hypothetica [SR:Synechocystis sp. (strain:PCC680 PCC6803 complete genome, 25/27, 3138 [RE:80528] [DI:complement]</pre>	B:pir2] l prote 3) DNA]	>gp:[in] [G [DB:g	GI:d1011491:g1001353] [LN:SYCSLLLH] N:clpP] [OR:Synechocystis sp.] enpept-bct1] [DE:Synechocystis sp.		
ORF Name	NTID	AAID	NT AA score probability		
AT7503000991_1445930_f2_152	1948	5720	621 206 103 0.0023		
Description					
	[GI:g29 OR:Aqui:	83776] fex ae	[LN:AE000736] [AC:AE000736:AE000657] plicus] [DB:genpept-bct2] [DE:Aquifex		
ORF Name	NTID	AAID	NT AA LengthLength score probability		
A17503000991_14460882_c1_443	1949	5721	1722 573 1304 4.9e-133		
<pre>gp:[GI:g1022726] [LN:SHU35635] [AC:U haemolyticus] [SR:Staphylococcus hae [DE:Staphylococcus haemolyticus IS12 [LE:1101] [RE:1922] [DI:complement]</pre>	molytic	us str	ain=Y176] [DB:genpept-bct1]		
ORF Name	NTID	AAID	NT AA score probability		
AT7503000991_14460887_£1_3	1950	5722	153 50 115 1.7e-06		
Description gp:[GI:g1022726] [LN:SHU35635] [AC:U haemolyticus] [SR:Staphylococcus hae [DE:Staphylococcus haemolyticus IS12 [LE:1101] [RE:1922] [DI:complement]	molytic	ıs str	ain=Y176] [DB:genpept-bct1]		
ORF Name	NTID	AAID	NT AA LengthLength		
AI7503000991_14484553_c2_467	1951	5723	1356 451 1726 9.3e-178		
Description sp:[LN:G6PA_BACST] [AC:P13375] [GN:PGIA] [OR:BACILLUS STEAROTHERMOPHILUS] [EC:5.3.1.9] [DE:ISOMERASE A)] [SP:P13375] [DB:swissprot] >pir:[LN:NUBSSA] [AC:S15936:S06196] [PN:glucose-6-phosphate isomerase, A:phosphoglucose isomerase:phosphohexose isomerase] [CL:glucose-6-phosphate isomerase] [OR:Bacillus stearothermophilus] [EC:5.3.1.9] [DB:pirl] >gp:[GI:g40046] [LN:BSPGIA] [AC:X16639] [OR:Bacillus stearothermophilus] [DB:genpept-bct1] [DE:Bacillus stearothermophilus pgiA gene for phosphoglucoisomeraseisoenzyme A (EC 5.3.1.9).] [NT:phosphoglucose isomerase A (AA 1-449)] [SP:P13375] [LE:95] [RE:1444] [DI:direct]					
ORF Name	NTID	AAID	NT AA LengthLength		
AI7503000991_14648512_c3_585	1952	5724	126 41		
<u>Description</u>	. 				
NO-HIT					

ORF Name	NTID	AAID	NT AA score probability
A17503000991_14656432_c3_582	1953	5725	
Description	L		
gp:[GI:g3158514] [LN:CELT04D1] [AC:A [DB:genpept-inv1] [DE:Caenorhabditis [LE:10425:10567:11420:12480] [RE:105	elegan	s cosm	id T04D1.]
ORF Name	NTID	AAID	NT AA score probability
AI7503000991_14884437_c1_437	1954	5726	165 54
Description			
NO-HIT			
ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000991_15038450_c2_469	1955	5727	135 44
Description			
NO-HIT			
ORF Name	NTID	AAID	NT AA score probability
A17503000991_16683437_f1_5	1956	5728	1023 340 571 2.3e-55
Description		•	
pir:[LN:A69863] [AC:A69863] [PN:con [OR:Bacillus subtilis] [DB:pir2] >gp [AC:Z99111:AL009126] [GN:ykrP] [FN:u [DE:Bacillus subtilis complete genom [NT:similar to hypothetical proteins	:[GI:e1: nknown] e (sect:	184942 [OR:B	:g2633723] [LN:BSUB0008] acillus subtilis] [DB:genpept-bct1] of 21): from 1394791to 1603020.]
ORF Name	NTID	AAID	NT AA score probability
A17503000991_16832562_f1_67	1957	5729	804 267 320 4.5e-36
Description			
<pre>pir:[LN:H69843] [AC:H69843] [PN:hyp subtilis] [DB:pir2] >gp:[GI:e1183175 [GN:yjbH] [FN:unknown] [OR:Bacillus complete genome (section 7 of 21): f [DI:complement]</pre>	:g263350 subtili:	09] [Li s] [DB	N:BSUB0007] [AC:Z99110:AL009126] :genpept-bct1] [DE:Bacillus subtilis
ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000991_187683_c1_417	1958	5730	228 75 256 1.3e-23
Description		· · · · · · · · · · · · · · · · · · ·	
sp:[LN:BIN3_STAAU] [AC:P20384] [GN:B DNA-INVERTASE BIN3 (TRANSPOSON TN552 [LN:SABINR3] [AC:X16298] [OR:Staphylococcus aureus plasmid pI transposon TN552.] [NT:bin3 product])] [SP:1 ococcus 9789 DN	P20384 aureus A with] [DB:swissprot] >gp:[GI:g398182] s] [DB:genpept-bct1] binR and bin3 genes,derived from
cramaposon insaz.1 [nr:bins product]	[35:52]	7704]	[mm.roas] [ve:res/] [hr:driece]

ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	score	probability
AI7503000991 195177 c2 521	1959	5731	141	46		2.6e-08
Description		L	لـــال		J L	
pir:[LN:S42239] [AC:S42239] [PN:hyp [DB:pir2] >gp:[GI:g501834] [LN:PNS1C from Staphylococcus aureus, plasmid Staphylococcus aureus) encodingtetra C; putative] [LE:1870] [RE:2409] [DI	G] [AC: pTP5 DN cycline	M16217 A] [DB -resis] [OR:I :genpep	Plasmid ot-bct1	l pNS1]	[SR:Plasmid pNS1 Plasmid pNS1 (from
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000991_19531626_f3_304	1960	5732	126	41]	
Description NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000991_19547938_c2_498	1961	5733	372	123	224	7.4e-18
Description			ــــــا كــــــــا		. —	
sp:[LN:G6PD_LEUME] [AC:P11411] [GN:Z [DE:GLUCOSE-6-PHOSPHATE 1-DEHYDROGEN						_
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000991_19566553_c1_428	1962	5734	159	52		
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000991_1960952_c2_473	1963	5735	906	301	575	8.7e-56
Description						
pir:[LN:C69837] [AC:C69837] [PN:5-o:yisK] [GN:yisK] [CL:2-hydroxyhepta-subtilis] [DB:pir2] >gp:[GI:e1183077 [GN:yisK] [FN:unknown] [OR:Bacillus complete genome (section 6 of 21): f:5-oxo-1,2,5-tricarboxilic-3-penten ac>gp:[GI:e1173509:g2145376] [LN:BSY09-[DB:genpept-bct1] [DE:B.subtilis 54k] with oxo-tricarboxilic-pentene] [LE:	2,4-die: :g26334 subtili rom 999 cid] [Li 476] [A	ne-1,7 11] [Li s] [DB 501 to: E:1522: C:Y094' ic DNA	-dioate N:BSUB0 :genpep 1209940 [RE 76] [PN fragme	isome 006] [ot-bct1] [NT ::15312 [:YisK]	rase] AC:Z99] [DE: :simil 1] [DI [OR:E NT:put	[OR:Bacillus 109:AL009126] Bacillus subtilis ar to :direct] acillus subtilis]
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000991_19610442_f1_56	1964	5736	246	81	219	4.6e-18
Description						
<pre>gp:[GI:d1045996:g5360820] [LN:D86934] aureus] [SR:Staphylococcus aureus (staphylococcus aureus [DB:genpept] [DE:Staphylococcus aureus [NT:ORF N026; putative] [LE:19527] []</pre>	train:N us gene:	315) DN s, mec	NA, clo region	ne_lib , part	:libra	ry of N31]

ORF Name	NTID	AAID	NT AA LengthLength	score	probability
AI7503000991_19790902_c2_493	1965	5737	813 270	919	3.1e-92
Description		<u> </u>			
pir:[LN:F69844] [AC:F69844] [PN:con [CL:conserved hypothetical protein E >gp:[GI:e1183181:g2633515] [LN:BSUBC [OR:Bacillus subtilis] [DB:genpept-b 7 of 21): from 1194391to 1411140.] [[RE:43541] [DI:direct]	HI0072] 0007] [A oct1] [D	OR:Ba C:Z991 E:Baci	cillus subtil 10:AL009126] llus subtilis	lis] [I [GN:y] s comp]	DB:pir2] jbN] [FN:unknown] Lete genome (section
ORF Name	NTID	AAID	NT AA LengthLength	score	probability
A17503000991_20517062_c2_478 Description	1966	5738	1002 333	961	1.1e-96
pir:[LN:F69842] [AC:F69842] [PN:3-c] [GN:yjaX] [CL:3-oxoacyl-[acyl-carri] [DB:pir2] >gp:[GI:el183136:g2633470] [FN:unknown] [OR:Bacillus subtilis] genome (section 6 of 21): from 99950 acyl-carrier protein] [LE:208189] [R[LN:BSUB0007] [AC:Z99110:AL009126] [DB:genpept-bct1] [DE:Bacillus subtili94391to 1411140.] [NT:similar to 3 [RE:14237] [DI:direct]	er-prot [LN:BS] [DB:gen 01 to120 EE:20912 [GN:yjaX]	ein] s UB0006 pept-b 9940.] 7] [DI] [FN: plete	ynthase III]] [AC:Z99109: ct1] [DE:Bac: [NT:similar :direct] >gp: unknown] [OR: genome (secti	[OR:Ba :AL0091 illus s to 3-c :[GI:e1 :Bacill	acillus subtilis] L26] [GN:yjaX] subtilis complete exoacyl- L183153:g2633487] Lus subtilis] of 21): from
			NT AA		
ORF Name	NTID	AAID	LengthLength	score	probability
A17503000991_20585302_f2_149	1967	5739	126 41	1	
Description NO-HIT				•	
ORF Name	NTID	AAID	NT AA LengthLength	score	probability
A17503000991_20739037_c1_452	1968	5740	426 141	690	5.7e-68
Description gp:[GI:g2267243] [LN:SEU71377] [AC:U [OR:Staphylococcus epidermidis] [DB: autolysin AtlE and putativetranscrip [LE:6867] [RE:7286] [DI:complement]	genpept	-bct1]	[DE:Staphylo	coccus	epidermidis
ORF Name	NTID	AAID	NT AA LengthLength	score	probability
AI7503000991_2125903_f3_313	1969	5741	1239 412	197	6.2e-13
Description				الـــــا ،	
pir:[LN:D71112] [AC:D71112] [PN:hyp [OR:Pyrococcus horikoshii] [DB:pir2] [AC:AP000003:AB009484:AB009485:AB009 hypothetical protein] [GN:PH0667] [O	>gp:[G 486:AB0	I:d103 09487:	0701:g3257075 AB009488:AB00	[LN:	AP000003] [PN:413aa long

(strain:OT3) DNA] [DB:genpept-bct1] [DE:Pyrococcus horikoshii OT3 genomic DNA, 544001-777000 nt. position(3/7).] [LE:51915] [RE:53156] [DI:complement]

ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	score	probability
A17503000991_21571937_£3_380	1970	5742	213	70	54	0.015
Description			JLJ		J L	
pir:[LN:F71120] [AC:F71120] [PN:hyp [OR:Pyrococcus horikoshii] [DB:pir2] [AC:AP000003:AB009484:AB009485:AB009 hypothetical protein] [GN:PH0733] [C (strain:OT3) DNA] [DB:genpept-bct1] 544001-777000 nt. position(3/7).] [L	>gp:[G 486:AB0 R:Pyroc [DE:Pyr	I:d103 09487: occus ococcu	0767:g3 AB00948 horikos s horik	257141 8:AB00 hii] (oshii	L] [LN 09489] [SR:Py: OT3 ge	:AP000003] [PN:150aa long rococcus horikoshii enomic DNA,
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
AI7503000991_21594202_f2_167 Description	1971	5743	1176	391	1674	3.0e-172
						caphylococcus genes.] [LE:5160]
ORF Name	NTID	AAID	<u>NT</u> Length!	<u>AA</u> Length	score	probability
AI7503000991_21753125_c2_481	1972	5744	1194	397	664	2.0e-79
Description	·				,	
pir:[LN:S16649] [AC:S16649:S18271:E6 (permease):dciAC protein] [GN:dppC:d [OR:Bacillus subtilis] [DB:pir2] >gp [PN:DppC] [GN:dppC] [FN:ABC-type dip [DB:genpept-bct1] [DE:Bacillus subti [LE:14301] [RE:15263] [DI:direct] >g [AC:Z99110:AL009126] [PN:dipeptide A subtilis] [DB:genpept-bct1] [DE:Baci from 1194391to 1411140.] [NT:alterna [DI:direct]	ciAC] :[GI:e1 eptide lis 168 p:[GI:e] BC trans	[CL:ol 181495 transpo 56 kb 118331 sporte: btilis	igopept :g26320 ort sys DNA fr 4:g2633 r (perm comple	ide pe 15] [I tem] [agment 648] [ease)] te gen	ermease IN:BSA OR:Bac Detwe LN:BSU [GN:C	e protein oppB] J2571] [AC:AJ002571] Cillus subtilis] een xlyA and ykoR.] JB0007] AppC] [OR:Bacillus section 7 of 21):
ORF Name	NTID	AAID	<u>NT</u> Lengthl	<u>AA</u> Length	score	probability
AI7503000991 22272583 f3 291	1973	5745		40		0.0040
Description			JL		L	
gp:[GI:g4098413] [LN:LOU77495] [AC:U [OR:Leuconostoc oenos bacteriophage bacteriophage 10MC putative integras recombinase] [LE:119] [RE:1165] [DI:	10MC] [] e (int)	DB:genp	pept-ph	g] [DE	:Leuco	nostoc oenos
ORF Name	NTID	AAID	<u>NT</u> LengthI	<u>AA</u> Length	score	probability
AI7503000991_22456512_f3_286	1974	5746	147	48		

Description NO-HIT

ORF Name	NTID	AAID	NT	<u>AA</u>	score	probability
AI7503000991 22462787 cl 392	<u> </u>	5747	Length	174		1.5e-51
Description			لـــــالـ			
sp:[LN:LEPH_STAAU] [AC:P72364] [GN:SPEPTIDASE IA] [SP:P72364] [DB:swissperset] [PN:type-I signal peptidase SpsA] [GE:Staphylococcus andtype-I signal peptidase SpsB (spspeptidase homologue; protein lacks]	orot] >gr GN:spsA] saureus GB) gene	o:[GI: [OR:S type- , comp	g159580 taphylo I signa lete co	09] [Li ococcus al pept ds.] [1	N:SAU69 s aureu tidase NT:inac	5000] [AC:U65000] us] SpsA (spsA) gene,
ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	score	probability
AI7503000991_22665887_c3_598	1976	5748	129	42		2.4e-07
Description		<u> </u>				
pir:[LN:D69838] [AC:D69838] [PN:con [OR:Bacillus subtilis] [DB:pir2] >gp [AC:Z99109:AL009126] [GN:yisU] [FN:u [DE:Bacillus subtilis complete genom [NT:similar to hypothetical proteins >gp:[GI:e1173520:g2145387] [LN:BSY09 [DB:genpept-bct1] [DE:B.subtilis 54k [RE:23869] [DI:complement]	o:[GI:ell unknown] ne (secti s] [LE:16 9476] [AC	183088 [OR:B ion 6 55981] C:Y094	:g26334 acillus of 21): [RE:16 76] [PN	122] [1 s subt: from 56643] U:YisU]	LN:BSUE ilis] 999501 [DI:co] [OR:E	B0006] [DB:genpept-bct1] L to1209940.] complement] Bacillus subtilis]
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000991_22703588_f2_137	1977	5749	954	317	803	6.0e-80
Description sp:[LN:MENA_BACSU] [AC:P39582] [GN:MENA_BACSU] [AC:P39582] [GN:MENA_BERNYLTRANSFERASE]] [SETAC:S39661:B70050] [PN:menaquinone [GN:ywaB] [CL:quinone biosynthesis page:[GI:g413930] [LN:BSGENR] [AC:X73 [DB:genpept-bct1] [DE:B.subtilis generates [RE:6839] [DI:direct] page:[GI:e11863 [GN:ywaB] [FN:unknown] [OR:Bacillus complete genome (section 20 of 21): ipa-6d; similar to quinone] [SP:P395]	biosynth homolog 124] [GN comic rec 48:g2636 subtilis	[DB nesis] ywaB] Y:ipa-(gion () 3384] [DB p8401te	swissgoroteing [OR:Based] [OR:Based] [OR:Based] [OR:BSU] [LN:BSU] [EN:BSU] [OR:BSU]	prot] : n homologicallus a:Bacil 333).] JB0020] ot-bct1 550.]	>pir:[I log ywa s subti llus su [SP:E [AC:Z L] [DE:	AN:S39661] AB:protein ipa-6d] AB:protein ipa-6d] Abtilis] AB:pir2] Abtilis] AB:pir2] AB:AB:AB:AB:AB:AB:AB:AB:AB:AB:AB:AB:AB:A
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000991_23438827_c3_566	1978	5750	129	42]	
Description NO-HIT						
110 1111		and the second s	ed Talantin (Talantin III) un colombusche		The second secon	
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000991_23535910_c1_391	1979	5751	603	200	·	9.8e-48
Description					. ———	
gp:[GI:g4185300] [LN:AF089862] [AC:A [DB:genpept-bct2] [DE:Staphylococcus						

464

andtype-I signal peptidase SipB (sipB) genes, complete cds; andunknown gene.] [NT:Orf1;

similar to Bacillus subtilis yhjE] [LE:271] [RE:948] [DI:direct]

ORF Name AI7503000991_23593932_t2_166 Description NO-HIT	NTID	<u>AAID</u> 5752	NT AA LengthLength	probability
ORF Name A17503000991_23611563_c3_600 Description NO-HIT	NTID 1981	<u>AAID</u> 5753	NT AA score LengthLength	probability
ORF Name A17503000991_23625637_c2_526 Description NO-HIT	NTID 1982	<u>AAID</u> 5754	NT AA score LengthLength	probability
ORF Name AI7503000991_23634641_c1_398 Description pir: [LN:G69842] [AC:G69842] [PN:3-o [CL:3-oxoacyl-[acyl-carrier-protein] synthase I homology] [OR:Bacillus su >gp:[GI:e1183137:g2633471] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-b 6 of 21): from 999501 to1209940.] [N [LE:209150] [RE:210391] [DI:direct] [AC:Z99110:AL009126] [GN:yjaY] [FN:u [DE:Bacillus subtilis complete genom [NT:similar to 3-oxoacyl- acyl-carri	syntha: btilis] 006] [Actl] [Di T:simil: >gp:[GI nknown] e (sect:	se I: [EC:2 C:Z991 E:Baci ar to :e1183 [OR:B ion 7	carrier-protein] syr 3-oxoacyl-[acyl-carr .3.1.41] [DB:pir2] 09:AL009126] [GN:yja 1lus subtilis comple 3-oxoacyl- acyl-carr 154:g2633488] [LN:BS acillus subtilis] [I of 21): from 1194391	rier-protein] aY] [FN:unknown] ete genome (section rier protein] SUB0007] DB:genpept-bct1]
ORF Name AI7503000991_23678800_c3_561 Description pir:[LN:H69841] [AC:H69841] [PN:con [CL:conserved hypothetical protein M >gp:[GI:e261991:g1620928] [LN:BS168N [OR:Bacillus subtilis] [DB:genpept-b [DI:direct] >gp:[GI:e1183118:g263345 [FN:unknown] [OR:Bacillus subtilis] genome (section 6 of 21): from 99950 [LE:192227] [RE:192535] [DI:direct] [AC:Y09476] [PN:YitW] [OR:Bacillus signomic DNA fragment.] [NT:putative	J1129] PRB] [AG ct1] [DI 2] [LN:I [DB:gen] 1 to1209 >gp:[GI ubtilis]	[OR:Ba C:Z795 E:B.su BSUB00 pept-b 9940.] :e1173	etical protein yitW] cillus subtilis] [DE 80] [GN:putative orf btilis nprB gene.] [06] [AC:Z99109:AL009 ct1] [DE:Bacillus su [NT:similar to hypo 550:g2145417] [LN:BS genpept-bct1] [DE:B.	3:pir2] [FN:unknown] [LE:6256] [RE:6564] [2126] [GN:yitW] [abtilis complete [5409476] [5409151 54kb
ORF Name A17503000991_2392300_t2_176 Description	NTID 1985	<u>AAID</u> 5757	NT AA score LengthLength 138 45	probability

ORF Name	MIID	<u> </u>	Length:	Length	20010	probabili	<u>L</u> Y
AI7503000991_24022191_f2_232	1986	5758	222	73] [97]	3.9e-05	
Description	I				·		
pir: [LN:G69854] [AC:G69854] [PN:hypsubtilis] [DB:pir2] >gp: [GI:e1183130] [GN:yjzD] [FN:unknown] [OR:Bacillus complete genome (section 6 of 21): f [DI:complement] >gp: [GI:e1183147:g26] [FN:unknown] [OR:Bacillus subtilis] genome (section 7 of 21): from 11943 >gp: [GI:d1023798:g2564027] [LN:D8637] subtilis (strain:CU741) DNA] [DB:gengenes, partial and complete cds.] [N [DI:complement]	:g263346 subtilis rom 9995 33481] [[DB:genp 91to 141 6] [AC:Do	4] [LM] [DB: 01 to: LN:BSU ept-bo 1140.] 86376]	S:BSUBO genpep 209940 JBOOO7] t1] [D [LE:9 [OR:B	006] [t-bct1 .] [LE [AC:Z E:Baci 807] [acillu lus su	[AC: Z99 .] [DE: 2: 20469 399110: .llus s [RE: 999 is subt	P109:AL00912 Bacillus su P7] [RE:2048 AL009126] [Subtilis comp P2] [DI:comp Pilis] [SR:B	btilis 82] GN:yjzD] splete lement] acillus
ORF Name	NTID	AAID	<u>NT</u> Length:	<u>AA</u> Length	score	probabili	ty
AI7503000991_24024142_c2_496	1987	5759	315	104	93	0.00010	· · ·
Description					· · · · · · · · · · · · · · · · · · ·		
gp:[GI:g2654481] [LN:BFU89914] [AC:U [OR:Bacillus firmus] [DB:genpept-bct protein, hypothetical 8.9 kDaprotein kDaprotein, putative thiosulfate sul transcriptional regulator and hypoth Na+/H+ antiporter homolog gene, part	2] [DE:Ba , hypothe furtrans: etical 1	acillu etical ferase 3.2 kI	s firm 10.1 , hypo a prot	us hyp kDa pr thetic eingen	otheti otein, al 16. es, co	cal 34.0 kD hypothetic 1kDa mplete cds,	al 21.0 and
ORF Name	NTID .	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probabili	ty
A17503000991_24070137_c3_602	1988	5760	150	49	Ì		
Description			[<u> </u>		•		
NO-HIT							
ORF Name	NTID	AAID	<u>NT</u> Length]	<u>AA</u> Length	score	probabili	ty
A17503000991_24105342_c1_448	1989	5761	831	276	1242	1.8e-126	
Description					_		
pir:[LN:F69656] [AC:F69656:A42715:S2 synthase:dihydroxynaphthoate synthas menB:menaquinone biosynthesis enzyme [CL:naphthoate synthase:enoyl-CoA hy [EC:4.1.3.36] [DB:pir2] >gp:[GI:e118 [PN:dihydroxynapthoic acid synthetas [OR:Bacillus subtilis] [DB:genpept-b genome (section 16 of 21): from 2997 [DI:complement] >gp:[GI:g2293148] [Lisynthase] [GN:menB] [OR:Bacillus subtrnB-dnaB genomic region.] [LE:29401	e: dihydr /enoyl Co dratase l 5953:g263 e] [GN:me ctl] [EC 771to 323 N:AF00822 tilis] [I	roxyna DA hyd nomolc 35564] enB] [:4.1.3 13410. DB:gen	pthoic ratase gy] [O [LN:B FN:men .36] [I] [LE: C:AF00 pept-b	acid homol R:Baci SUB001 aquino DE:Bac 150211 8220] ct2] [synthe og] [G llus s 6] [AC ne bio illus] [RE: [PN:di	tase N:menB] ubtilis] :Z99119:AL0 synthesis] subtilis co 151026] hydroxynaph	mplete thoate
ORF Name	NTID 2	AAID	<u>NT</u> Length I	<u>AA</u> Length	score	probabili	EY
AI7503000991_24225010_c2_536	1990	5762	168	55			
Description NO-HIT	L		(

ORF Name	NTID AAID LengthLength score probability
AI7503000991_24270451_c1_432	1991 5763 135 44
Description NO-HIT	
ORF Name	NTID AAID NT AA score probability
A17503000991_24409428_c1_438	1992 5764 258 85 357 1.1e-32
[OR:Staphylococcus aureus] [DB:genper	[38429] [PN:recombination protein] [GN:pre] [pt-bctl] [DE:Staphylococcus aureus chloramphenicol [uence.] [LE:2074] [RE:3285] [DI:direct]
ORF Name	NTID AAID NT AA score probability
A17503000991_24412517_c3_581	1993 5765 393 130 213 1.2e-16
[PN:probable glucose-6-phosphate 1-de [CL:glucose-6-phosphate dehydrogenase >gp:[GI:d1013296:g1303961] [LN:BACJH6 subtilis] [SR:Bacillus subtilis (stra [DE:Bacillus subtilis DNA, 283 Kb reg [RE:240494] [DI:complement] >gp:[GI:e [AC:Z99116:AL009126] [GN:yqjJ] [FN:um [DE:Bacillus subtilis complete genome	e] [OR:Bacillus subtilis] [EC:1.1.1.49] [DB:pir2] [642] [AC:D84432:D82370] [PN:YqjJ] [OR:Bacillus ain:JH642(trpC2 PheA1)) DNA] [DB:genpept-bct1] gion containing skin element.] [LE:239025]
ORF Name	NTID AAID NT AA score probability
[DE:SIGNAL PEPTIDASE IB, (SPASE IB) >gp:[GI:g1595810] [LN:SAU65000] [AC:U] [OR:Staphylococcus aureus] [DB:genper] peptidase SpsA (spsA) gene, andtype-1	SB] [OR:STAPHYLOCOCCUS AUREUS] [EC:3.4.21.89] (LEADER PEPTIDASE IB)] [SP:P72365] [DB:swissprot] U65000] [PN:type-I signal peptidase SpsB] [GN:spsB] pt-bct2] [DE:Staphylococcus aureus type-I signal I signal peptidase SpsB (spsB) gene, complete cds.] se, serine] [LE:580] [RE:1155] [DI:direct]
ORF Name	NTID AAID NT AA probability LengthLength
<pre>epidermidis] [DB:genpept-bct1] [DE:St</pre>	1995 [5767 4008 1335 7001 0.0 71377] [PN:autolysin AtlE] [OR:Staphylococcus taphylococcus epidermidis autolysin AtlE and lR genes, complete cds.] [NT:primary attachment to
ORF Name A17503000991 24428543 £3 297	NTID AAID NT AA score probability

AΑ NT ORF Name NTID AAID score probability LengthLength A17503000991 24475252 £3 289 1997 5769 159 Description NO-HIT NT AΑ ORF Name \mathtt{NTID} AAID score probability LengthLength A17503000991_24646886_c3_594 5770 1557 1998 518 2.0e-23 Description pir:[LN:B71973] [AC:B71973] [PN:2',3'-cyclic-nucleotide 2'-phosphodiesterase] [GN:cpdB [CL:2',3'-cyclic-nucleotide 2'-phosphodiesterase: 2',3'-cyclic-nucleotide 2'-phosphodiesterase homology:phosphoesterase core homology] [OR:Helicobacter pylori] [SR:strain J99, , strain J99] [SR:strain J99,] [DB:pir2] >gp:[GI:g4154615] [LN:AE001449] [AC:AE001449:AE001439] [PN:2',3'-CYCLIC-NUCLEOTIDE 2'-PHOSPHODIESTERASE] [GN:cpdB] [OR:Helicobacter pylori J99] [DB:genpept-bct2] [DE:Helicobacter pylori, strain J99 section 10 of 132 of the completegenome.] [NT:similar to H. pylori 26695 gene HP0104] [LE:139] [RE:1884] [DI:complement] NT AA ORF Name NTID AAID score LengthLength A17503000991 24648412 cl 406 1999 642 3.1e-67 Description pir:[LN:E69844] [AC:E69844] [PN:GTP pyrophosphokinase homolog yjbM] [GN:yjbM] [CL:GTP pyrophosphokinase related protein] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1183180:g2633514] [LN:BSUB0007] [AC:Z99110:AL009126] [GN:yjbM] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 7 of 21): from 1194391to 1411140.] [NT:similar to GTP pyrophosphokinase] [LE:42087] [RE:42722] [DI:direct] NT AΑ ORF Name NTID AAID score probability LengthLength AI7503000991 24745437_c3_539 2000 1263 420 1659 1.2e-170 Description sp:[LN:YPCA BACSU] [AC:P50735] [GN:YPCA] [OR:BACILLUS SUBTILIS] [EC:1.4.1.-] [DE:(EC 1.4.1.-)] [SP:P50735] [DB:swissprot] >pir:[LN:G69933] [AC:G69933] [PN:glutamate dehydrogenase homolog ypcA] [GN:ypcA] [CL:glutamate dehydrogenase (NAD(P)+)] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1183741:q2634714] [LN:BSUB0012] [AC:Z99115:AL009126] [GN:ypcA] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 12 of 21): from 2195541to 2409220.] [NT:similar to glutamate dehydrogenase] [SP:P50735] [LE:205764] [RE:207044] [DI:complement] >gp:[GI:e1185565:g2634731] [LN:BSUB0013] [AC:Z99116:AL009126] [GN:ypcA] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 13 of 21): from 2395261to 2613730.] [NT:similar to qlutamate dehydrogenase] [SP:P50735] [LE:6044] [RE:7324] [DI:complement] >gp:[GI:g1146206] [LN:BACSERA] [AC:L47648] [PN:glutamate dehydrogenase] [GN:ypcA] [OR:Bacillus subtilis] [DB:genpept-bct2] [DE:Bacillus subtilis phosphoglycerate dehydrogenase (serA), ypaA, ferredoxin (fer), ypbB, recS, ypbD, ypbE, ypbF, ypbG, ypbH, glutamate dehydrogenase (ypcA), ypdA, ypdB, ypdC, spore cortexlytic enzyme (sleB), ypeB, ypfA, ypfB, cytidine

dependentglycerol-3-phosphate dehydrogenase (glyc), yphE and yphF genes,complete cds.]

[NT:similar to Clostridium difficile glutamate] [LE:9353] [RE:10633] [DI:direct]

monophosphatekinase (cmk), ypfD, ypgA, yphA, yphB, yphC, NAD+

ORF Name	NTID	AAID	NT AA LengthLength score probability
AI7503000991_24804677_c3_597	2001	5773	366 121 117 1.5e-06
Description gp:[GI:g2541936] [LN:PSU27310] [AC:U [DB:genpept-bct2] [DE:Pseudomonas sy sequence.] [NT:ORF6; similar to Pseu [DI:direct]	ringae	phaseo	lotoxin gene cluster, complete
ORF Name	NTID	AAID	NT AA core probability
AI7503000991_25392826_f2_175	2002	5774	1146 381 557 7.0e-54
Description			
pir:[LN:G70007] [AC:G70007] [PN:con [CL:Bacillus subtilis conserved hypo [DB:pir2] >gp:[GI:e1184257:g2635675] [FN:unknown] [OR:Bacillus subtilis] genome (section 17 of 21): from 3197 proteins] [LE:64687] [RE:65796] [DI:	thetica [LN:BS [DB:gen 001to 3	l prot UB0017 pept-b 414420	ein yueF] [OR:Bacillus subtilis]] [AC:Z99120:AL009126] [GN:yueF] ct1] [DE:Bacillus subtilis complete
ORF Name	NTID	AAID	NT AA score probability
A17503000991_25429512_c3_593	2003	5775	1392 463 1130 1.3e-114
[CL:Na+-ATP synthase chain J] [OR:Ba >gp:[GI:e1184940:g2633721] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-b 8 of 21): from 1394791to 1603020.] [LE:20743] [RE:22092] [DI:direct]	008] [A	C:Z991 E:Baci	11:AL009126] [GN:ykrM] [FN:unknown] llus subtilis complete genome (section
ORF Name	NTID	AAID	NT AA score probability
A17503000991_25430316_c2_522	2004	5776	171 56 171 5.6e-13
Description pir:[LN:QQSA8T] [AC:A04493] [PN:hyp [DB:pir1]	othetica	al pro	tein B-189] [OR:Staphylococcus aureus]
ORF Name	NTID	AAID	NT AA score probability
A17503000991_26178400_c3_542	2005	5777	186 61 71 0.042
Description sp:[LN:AOPP_HUMAN] [AC:P30044] [OR:H [DB:swissprot]	OMO SAP	IENS]	[SR:, HUMAN] [DE:71B)] [SP:P30044]
ORF Name	NTID	AAID	NT AA score probability
AI7503000991_26212787_£3_335	2006	5778	612 203 308 1.7e-27
Description			
<pre>pir:[LN:C69844] [AC:C69844] [PN:hyp- subtilis] [DB:pir2] >gp:[GI:e1183178 [GN:yjbK] [FN:unknown] [OR:Bacillus complete genome (section 7 of 21): f [DI:complement]</pre>	:g26335: subtili:	12] [Li s] [DB	N:BSUB0007] [AC:Z99110:AL009126] :genpept-bctl] [DE:Bacillus subtilis

ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	score	probability
A17503000991_26377340_c2_488	2007	5779	999	332	208	4.6e-15
Description gp:[GI:e257629:g1771202] [LN:LLLVSFP] [OR:Lactococcus lactis] [DB:genpept-	bct1] [1	DE:L.1	actis O	RF, ge	enes ho	omologous to vsf-1
and pepF2 and gene encodingprotein he with vsf-1 gene (X73635)] [LE:934] [ransfe	erase.]	[NT:weak homology
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000991_26564012_c2_477	2008	5780	2616	871	2649	1.5e-275
Description gp:[GI:g3150046] [LN:AF016634] [AC:A [OR:Lactococcus lactis subsp. cremoric cremoris ClpB chaperone homolog (clpb (pur5) genes, completecds; and phosph gene, partial cds.] [LE:183] [RE:2786]	is] [DB B) andpl horibosy	:genpe nospho ylglyc	pt-bct2 ribosyl	[DE: formyl	Lactor glycin	occus lactis amide cyclo-ligase
ORF Name	NTID	AAID	NT Length	AA Tonath	score	probability
AI7503000991 26567557 c3 558	2009	5781		45	1	
Description					J	
NO-HIT						
ORF Name	NTID	AAID	NT Length:	<u>AA</u> Length	score	probability
A17503000991_26604662_c2_495	2010	5782		209]	
Description					J	
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000991_26775637_c1_447	2011	5783	1389	462	599	2.5e-58
Description sp:[LN:MENF_BACSU] [AC:P23973:P23972] [GN:MENF:ICSM] [OR:BACILLUS SUBTILIS] [EC:5.4.99.6] [DE:MENAQUINONE-SPECIFIC ISOCHORISMATE SYNTHASE,] [SP:P23973:P23972] [DB:swissprot] >pir:[LN:A69657] [AC:A69657:S27507:S27508:I39883] [PN:probable isochorismate synthase, menaquinone-specific menF] [GN:menF] [CL:isochorismate synthase] [OR:Bacillus subtilis] [EC:5.4.99.6] [DB:pir2] >gp:[GI:e1185956:g2635567] [LN:BSUB0016] [AC:Z99119:AL009126] [PN:menaquinone-specific isochorismate synthase] [GN:menF] [FN:menaquinone biosynthesis] [OR:Bacillus subtilis] [DB:genpept-bct1] [EC:5.4.99.6] [DE:Bacillus subtilis complete genome (section 16 of 21): from 2997771to 3213410.] [NT:alternate gene name: icsM, entC] [SP:P23973] [LE:153612] [RE:155027] [DI:complement] >gp:[GI:g2293145] [LN:AF008220] [AC:AF008220] [PN:isochorismate synthase] [GN:menF] [OR:Bacillus subtilis] [DB:genpept-bct2] [DE:Bacillus subtilis rrnB-dnaB genomic region.] [LE:25400] [RE:26815] [DI:direct] >gp:[GI:g1185288] [LN:BACMENAQOP] [AC:M74538:M74182:M74183] [PN:isochorismate synthase] [GN:menF] [OR:Bacillus subtilis] [SR:Bacillus subtilis (strain RB1) DNA] [DB:genpept-bct2] [DE:Bacillus subtilis menaquinone operon: menF, menD, menB and menEgenes, complete cds.] [NT:based on similarity to E. coli EntC, A. hydrophila] [LE:143] [RE:1558] [DI:direct]						
ORF Name	NTID	AAID	NT Lengthl	<u>AA</u> Length	<u>score</u>	probability
A17503000991_2848308_c3_607	2012	5784	135	44		
Description NO-HIT						

ORF Name	NTID	AAID	NT AA score probability
AI7503000991_289802_£2_227	2013	5785	
Description	l —————		الــــــــــــــــــــــــــــــــــــ
pir:[LN:F22845] [AC:F22845] [PN:hyp	pothetica	al pro	tein 6] [OR:mitochondrion Trypanosoma
brucei] [DB:pir2]		_	
ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000991_2922260_c3_574	2014	5786	384 127 174 2.7e-13
Description			
pir:[LN:D69844] [AC:D69844] [PN:hypsubtilis] [DB:pir2] >gp:[GI:e1183179 [GN:yjbL] [FN:unknown] [OR:Bacillus complete genome (section 7 of 21): f[DI:direct]	g263351: subtilis	L3] [L s] [DB	N:BSUB0007] [AC:Z99110:AL009126] :genpept-bct1] [DE:Bacillus subtilis
ORF Name	NTID	AAID	NT AA score probability
A17503000991_29335888_c3_562	2015	5787	135 44
Description			<u> </u>
NO-HIT			
ORF Name	NTID	AAID	NT AA LengthLength score probability
AI7503000991_29570302_f1_24	2016	5788	132 43
Description			
NO-HIT	<u> </u>		
ORF Name	NTID	AAID	NT AA score probability
A17503000991_31281253_c3_538	2017	5789	1221 406 1513 3.5e-155
Description		-	
sp:[LN:OAT_BACSU] [AC:P38021] [GN:RC			
[DE:AMINOTRANSFERASE]] [SP:P38021]			<pre>>pir:[LN:S55793] oxo-acid transaminase, rocD:ornithine</pre>
aminotransferase:ornithineoxo-acid			
[CL:ornithineoxo-acid aminotransfe	erase] [C	R:Bac	illus subtilis] [EC:2.6.1.13]
[DB:pir2] >gp:[GI:d1011955:g1064807]			
			[SR:Bacillus subtilis (strain:168)
DNA] [DB:genpept-bct1] [DE:Bacillus genesencoding 34 ORFs.] [LE:25845] [
[LN:BSROCDEF] [AC:X81802] [PN:ornith			
subtilis] [DB:genpept-bct1] [DE:B.su			
[LE:241] [RE:1446] [DI:direct] >gp:[[AC:Z99124:AL009126] [PN:ornithine a			
ornithine utilization] [OR:Bacillus			
[DE:Bacillus subtilis complete genom			
[SP:P38021] [LE:144229] [RE:145434]	[DI:comp	lemen	=]
ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000991_31300807_c1_436	2018	5790	
Description	L		·
NO-HIT			

ORF Name	NTID	AAID NT AA score probability				
AI7503000991_31525260_c3_579	2019	5791 135 44				
Description						
NO-HIT						
ORF Name	NTID	AAID Longth longth score probability				
AI7503000991 32062875 cl 439		<u> </u>				
	2020	5792 291 96 333 3.8e-30				
Description gp:[GI:g1731452] [LN:SAU38429] [AC:U [OR:Staphylococcus aureus] [DB:genpe resistance plasmid pKH7,complete seq	pt-bct1]	[DE:Staphylococcus aureus chloramphenicol				
ORF Name	NTID	AAID NT AA score probability				
AI7503000991_32067937_f2_247	2021	5793 1365 454 1790 1.5e-184				
Description						
[FN:pyridine nucleotide disulfide ox	idoreduo aureus	coenzyme A disulfide reductase gene,				
ORF Name	NTID	AAID NT AA score probability				
AI7503000991 32132183 c1 455	2022	5794 132 43				
Description						
NO-HIT						
ORF Name	NTID	AAID NT AA score probability				
A17503000991 32457312 c2 501	2023	[5795 [495 164 233 1.5e-19				
Description						
gp:[GI:e303881:g1850807] [LN:CPCPEAA] [AC:X71844] [PN:putative transposase] [OR:Clostridium perfringens] [DB:genpept-bct1] [DE:C.perfringens uapC, cpe, and nadC genes.] [LE:2477] [RE:2932] [DI:direct]						
ORF Name	NTID	AAID NT AA score probability				
AI7503000991_32635937_c1_429	2024	5796 1503 500 2239 4.1e-232				
Description						
	ccus aur	Y14370] [PN:UDP-N-acetylmuramyl-tripeptide ceus] [DB:genpept-bct1] [DE:Staphylococcus E:4722] [DI:complement]				
ORF Name	NTID	AAID NT AA score probability				
A17503000991_33364067_c1_431	2025	5797 1782 593 542 2.7e-52				
Description						
[OR:Streptococcus pneumoniae] [DB:gen	npept-bo putative ubunit o					

ORF Name	NTID AAID LengthLength score probability
AI7503000991_3371067_c2_482	2026 5798 954 317 1033 2.5e-104
protein oppF:sporulation initiation membrane protein malK:ATP-binding ca >gp:[GI:g143608] [LN:BACSPOOK] [AC:M [OR:Bacillus subtilis] [SR:B.subtili	59669] [PN:oligopeptide transport ATP-binding protein spo0KE] [GN:spo0KE:oppF] [CL:inner assette homology] [OR:Bacillus subtilis] [DB:pir2] M57689] [PN:sporulation protein] [GN:spo0KE] is (strain JH642) DNA, clones pDR20/21, pJL2/3 and a subtilis spo0K operon.] [LE:5152] [RE:6078]
ORF Name A17503000991_33710968_f2_124	NTID AAID NT AA score probability 2027 5799 453 150 742 1.7e-73
[DB:genpept-bct1] [DE:Staphylococcus	J71377] [OR:Staphylococcus epidermidis] s epidermidis autolysin AtlE and clR genes, complete cds.] [NT:ORF3] [LE:1957]
ORF Name A17503000991_3394540_f2_138 Description NO-HIT	NTID AAID NT AA score probability 2028 5800 312 103
ORF Name A17503000991_34006561_f3_312 Description NO-HIT	NTID AAID NT AA score probability 2029 5801 150 49
ENZYME (MOBILIZATION PROTEIN)] [SP:PG [AC:J01764:J01765] [OR:Plasmid pT181]	NTID AAID NT AA LengthLength score probability 2030 5802 273 90 290 1.4e-25 STAPHYLOCOCCUS AUREUS] [DE:PLASMID RECOMBINATION 203864] [DB:swissprot] >gp:[GI:g151683] [LN:PT1CG] L] [SR:Plasmid pT181 DNA from Staphylococcus aureus] complete genome.] [NT:Pre protein (plasmid [DI:direct]
	NTID AAID NT AA LengthLength score probability 2031 5803 1545 514 2291 1.3e-237 370] [AC:Y14370] [PN:peptide chain release factor 3] [DB:genpept-bct1] [DE:Staphylococcus aureus RF3, 20] [DI:complement]

ORF Name	NULLD	7 7 TD	NT AA GGORG Puchchilita
	NTID	AAID	Length Length score probability
A17503000991_34163562_f3_322	2032	5804	486 161 329 1.0e-29
Description pir: [LN:D69583] [AC:D69583] [PN:alc dehydrogenase:long-chain alcohol deh [EC:1.1.1.1] [DB:pir2] >gp:[GI:e2098 [PN:NAD alcohol dehydrogenase] [GN:a [DE:B.subtilis 23.9kb fragment from [LE:13725] [RE:14861] [DI:complement [AC:Z99117:AL009126] [PN:alcohol deh [DB:genpept-bct1] [DE:Bacillus subti 2599451to 2812870.] [LE:153270] [RE:	ydrogen 90:g210 dhB] [0 map pos] >gp:[ydrogen lis com	ase home ase home ase home ase] [mology] [OR:Bacillus subtilis] [LN:BS233DEG] [AC:X92868:X79978] llus subtilis] [DB:genpept-bct1] 233 degrees on thechromosome.] 83926:g2635142] [LN:BSUB0014] GN:adhB] [OR:Bacillus subtilis] genome (section 14 of 21): from
ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000991_34179031_f2_122	2033	5805	135 44
Description			
NO-HIT		A	
ORF Name	NTID	AAID	NT AA score probability
AI7503000991_34398505_£3_354	2034	5806	222 73
Description			J
NO-HIT			
ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000991_34407888_c3_572	2035	5807	729 242 288 1.9e-37
Description			
<pre>gp:[GI:g2952027] [LN:BFU88888] [AC:U firmus] [DB:genpept-bct2] [DE:Bacill synthase (cls)genes, complete cds.]</pre>	us firm	ıs Meci	A homolog (mecA) and cardiolipin
ORF Name	NTID	AAID	NT AA LengthLength score probability
AI7503000991_35158177_c1_408	2036	5808	1407 468 903 1.5e-90
	(strain	C-125	374] [PN:Ykok] [GN:ykok] [OR:Bacillus) DNA] [DB:genpept-bct1] [DE:Bacillus partialand complete cds.] [LE:1793]
ORF Name	NTID	AAID	NT AA score probability
A17503000991_35937827_c2_505	2037	5809	
Description			JL
pir:[LN:G69846] [AC:G69846] [PN:hypsubtilis] [DB:pir2] >gp:[GI:e1183205 [GN:yjcG] [FN:unknown] [OR:Bacillus complete genome (section 7 of 21): f.[DI:complement]	:g263353 subtilia	39] [L1 3] [DB	N:BSUB0007] [AC:Z99110:AL009126] :genpept-bct1] [DE:Bacillus subtilis

OPE Name	NULLD	NT AA GGOTO THOUSEN
ORF Name	NTID	AAID Length Length score probability
A17503000991_36128387_c3_621	2038	5810 165 55
Description		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
AI7503000991_36134717_c1_401	2039	
Description		
[GN:traC] [CL:dipeptide transport p >gp:[GI:g388269] [LN:AD1TRAC] [AC:L1 DNA] [DB:genpept-bct1] [DE:Plasmid p	rotein] .9532] [AD1 (fr	cAD1 binding protein precursor:TraC]] [OR:Enterococcus faecalis] [DB:pir2] [GN:traC] [OR:Plasmid pAD1] [SR:Plasmid pAD1 rom Enterococcus faecalis strain: DS16) cds, traA and traB genes, 3' end.] [LE:166]
ORF Name	NTID	AAID NT AA score probability
A17503000991_36229625_£3_295	2040	5812 204 67 195 1.6e-15
Description		
<pre>subtilis] [DB:pir2] >gp:[GI:e1184971 [GN:ykvS] [FN:unknown] [OR:Bacillus</pre>	:g26337 subtili	cal protein ykvS] [GN:ykvS] [OR:Bacillus 752] [LN:BSUB0008] [AC:Z99111:AL009126] is] [DB:genpept-bct1] [DE:Bacillus subtilis 94791to 1603020.] [LE:52338] [RE:52769]
ORF Name	NTID	AAID <u>NT AA</u> score probability
A17503000991_36611062_c3_557	2041	5813 420 139 174 2.7e-13
<pre>subtilis] [DB:pir2] >gp:[GI:e1183078 [GN:yisL] [FN:unknown] [OR:Bacillus complete genome (section 6 of 21): f [DI:direct] >gp:[GI:e1173510:g214537</pre>	:g26334 subtili: rom 999 7] [LN:	cal protein yisL] [GN:yisL] [OR:Bacillus 412] [LN:BSUB0006] [AC:Z99109:AL009126] is] [DB:genpept-bct1] [DE:Bacillus subtilis 9501 to1209940.] [LE:153237] [RE:153593] :BSY09476] [AC:Y09476] [PN:YisL] [OR:Bacillus 54kb genomic DNA fragment.] [NT:putative]
ORF Name AI7503000991_37777_c1_434 Description	NTID 2042	AAID NT AA score probability [5814] [168] [55]

NTAΑ ORF Name NTID probability LengthLength A17503000991 3954718 cl 394 3489 2043 5815 1162 6.7e-136 Description sp:[LN:ADDB BACSU] [AC:P23477] [GN:ADDB] [OR:BACILLUS SUBTILIS] [DE:ATP-DEPENDENT NUCLEASE SUBUNIT B] [SP:P23477] [DB:swissprot] >pir:[LN:A39432] [AC:A39432:S61272:A69583] [PN:ATP-dependent deoxyribonuclease chain B addB:ATP-dependent exonuclease synthesis protein AddB] [GN:addB] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:g142439] [LN:BACADDAA] [AC:M63489] [PN:ATP-dependent nuclease] [GN:addA] [OR:Bacillus subtilis] [SR:B.subtilis DNA] [DB:genpept-bct1] [DE:Bacillus subtilis ATP-dependent nuclease (addA) and (addB), andopen reading frame 3, partial cds.] [LE:502] [RE:4002] [DI:direct] >gp:[GI:e1183064:g2633398] [LN:BSUB0006] [AC:Z99109:AL009126] [PN:ATP-dependent deoxyribonuclease (subunit B)] [GN:addB] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 6 of 21): from 999501 to1209940.] [SP:P23477] [LE:136293] [RE:139793] [DI:direct] >gp:[GI:e1364880:g2226191] [LN:BSY14081] [AC:Y14081] [PN:ATP-dependent nuclease] [GN:addB] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis chromosomal DNA, region 92 degrees: regionbetween comK and addAB.] [NT:TTG start; see ref [3]; In EMBL entry M63489 this] [LE:18588] [RE:22088] [DI:direct] NTAΑ ORF Name NTID AAID score probability LengthLength A17503000991_4019193_c1_418 2044 5816 783 260 749 Description pir:[LN:G69845] [AC:G69845] [PN:enoyl-[acyl-carrier-protein] reductase (NADH), yjbW] [GN:yjbW] [CL:enoyl-[acyl-carrier-protein] reductase (NADH): short-chain alcohol dehydrogenase homology] [OR:Bacillus subtilis] [EC:1.3.1.9] [DB:pir2] >gp:[GI:e1183192:g2633526] [LN:BSUB0007] [AC:Z99110:AL009126] [GN:yjbW] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 7 of 21): from 1194391to 1411140.] [NT:similar to enoyl- acyl-carrier protein reductase] [LE:52836] [RE:53645] [DI:direct] NT AA ORF Name NTID AAID score probability Length Length AI7503000991 4070151 c2 472 2045 5817 447 148 165 2.4e-12 Description gp:[GI:e1173495:g2145362] [LN:BSY09476] [AC:Y09476] [PN:AddA] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:B.subtilis 54kb genomic DNA fragment.] [SP:P23478] [LE:<1] [RE:702] [DI:direct] NT AΑ ORF Name NTID AAID score probability LengthLength A17503000991_4080443_c1_409 2046 5818 1854 617 1.2e-161

Description

pir: [LN:A69845] [AC:A69845] [PN:Na+/H+ antiporter homolog yjbQ] [GN:yjbQ]

[OR:Bacillus subtilis] [DB:pir2] >qp:[GI:e1183184:q2633518] [LN:BSUB0007]

[AC:Z99110:AL009126] [GN:yjbQ] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1]

[DE:Bacillus subtilis complete genome (section 7 of 21): from 1194391to 1411140.]

[NT:similar to Na+/H+ antiporter] [LE:45437] [RE:47281] [DI:direct]

NT ORF Name NTID AAID score probability LengthLength A17503000991 4088962 c2 508 2047 5819 0.0065 Description pir: [LN:A70008] [AC:A70008] [PN:hypothetical protein yueH] [GN:yueH] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1184255:q2635673] [LN:BSUB0017] [AC:Z99120:AL009126] [GN:yueH] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 17 of 21): from 3197001to 3414420.] [LE:64085] [RE:64333] [DI:complement] NT AA ORF Name NTID AAID score probability LengthLength A17503000991 4094433 f1 72 5820 2048 999 332 1184 2.5e-120 Description sp:[LN:SYW BACSU] [AC:P21656] [GN:TRPS] [OR:BACILLUS SUBTILIS] [EC:6.1.1.2] [DE:(TRPRS)] [SP:P21656] [DB:swissprot] >pir:[LN:YWBS] [AC:JT0481:A32452:E69726] [PN:tryptophan--tRNA ligase,:tryptophanyl-tRNA synthetase] [GN:trpS] [CL:tryptophan--tRNA ligase] [OR:Bacillus subtilis] [EC:6.1.1.2] [DB:pir1] >gp:[GI:g143786] [LN:BACTRPSA] [AC:M24068] [OR:Bacillus subtilis] [SR:B.subtilis (strain QB928) DNA, clone pTSQ2] [DB:genpept-bct1] [DE:B.subtilis trpS gene encoding tryptophanyl-tRNA synthetase, complete cds.] [NT:tryptophanyl-tRNA synthetase (EC 6.1.1.2)] [LE:171] [RE:1163] [DI:direct] >qp:[GI:e1183162:q2633496] [LN:BSUB0007] [AC:Z99110:AL009126] [PN:tryptophanyl-tRNA synthetase] [GN:trpS] [OR:Bacillus subtilis] [DB:genpept-bct1] [EC:6.1.1.2] [DE:Bacillus subtilis complete genome (section 7 of 21): from 1194391to 1411140.] [SP:P21656] [LE:23195] [RE:24187] [DI:complement] NT ORF Name NTID AAID score probability LengthLength A17503000991_40957_c1_407 2049 858 285 3.2e-49 Description sp:[LN:YJBO BACSU] [AC:031613] [GN:YJBO] [OR:BACILLUS SUBTILIS] [DE:HYPOTHETICAL 31.5 KD PROTEIN IN MECA-TENA INTERGENIC REGION] [SP:031613] [DB:swissprot] >pir:[LN:G69844] [AC:G69844] [PN:conserved hypothetical protein yjb0] [GN:yjb0] [CL:conserved hypothetical protein HI0176] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1183182:g2633516] [LN:BSUB0007] [AC:Z99110:AL009126] [GN:yjb0] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 7 of 21): from 1194391to 1411140.] [NT:similar to hypothetical proteins] [SP:O31613] [LE:43604] [RE:44455] [DI:direct] NTAA ORF Name NTID AAID probability score LengthLength A17503000991 4100393 f3 379 2050 1215 404 Description pir: [LN:B69589] [AC:B69589] [PN:argininosuccinate synthase argG] [GN:argG] [CL:argininosuccinate synthase] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1184194:g2635410] [LN:BSUB0015] [AC:Z99118:AL009126] [PN:argininosuccinate synthase] [GN:argG] [FN:arginine biosynthesis] [OR:Bacillus subtilis] [DB:genpept-bct1] [EC:6.3.4.5] [DE:Bacillus subtilis complete genome (section 15 of 21): from 2795131to 3013540.] [LE:217075] [RE:218286] [DI:complement] >qp:[GI:e1185818:q2635429] [LN:BSUB0016] [AC:Z99119:AL009126] [PN:argininosuccinate synthase] [GN:argG] [FN:arginine biosynthesis] [OR:Bacillus subtilis] [DB:genpept-bct1] [EC:6.3.4.5] [DE:Bacillus subtilis complete genome (section 16 of 21): from 2997771to 3213410.] [LE:14435] [RE:15646] [DI:complement] >gp:[GI:g2293242] [LN:AF008220] [AC:AF008220] [PN:arginine succinate synthase] [GN:argG] [OR:Bacillus subtilis] [DB:genpept-bct2]

[DE:Bacillus subtilis rrnB-dnaB genomic region.] [LE:164781] [RE:165992] [DI:direct]

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000991_4173410_c2_504	2051	5823	792	263	485	3.0e-46
Description					- I	
pir:[LN:H69846] [AC:H69846] [PN:hypsubtilis] [DB:pir2] >gp:[GI:e1183206 [GN:yjcH] [FN:unknown] [OR:Bacillus complete genome (section 7 of 21): f[DI:complement]	:g26335 subtili	40] [L s] [DB	N:BSUB(:genper	007] ot-bct1	[AC:Z99 L] [DE:	9110:AL009126] :Bacillus subtilis
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
AI7503000991_4181312_c2_527	2052	5824	606	201	93	0.010
Description						
pir: [LN:A71455] [AC:A71455] [PN:pro [OR:Pyrococcus horikoshii] [DB:pir2] [AC:AP000001:AB009465:AB009464:AB009 hypothetical acetyltransferase] [GN:horikoshii (strain:OT3) DNA] [DB:gen DNA, 1-287000 nt. position (1/7).] [in] [LE:262066] [RE:262584] [DI:comp	>gp:[G 466:AB0 PH0296] pept-bc NT:simi	I:d103 09467: [OR:P t1] [D	0311:g3 AB00946 yrococo E:Pyroc	256685 8:AB00 cus hor coccus	[LN: 9469] rikoshi horiko	AP000001] [PN:172aa long ii] [SR:Pyrococcus oshii OT3 genomic
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
AI7503000991_4335752_f1_4	2053	5825	171	56]	
Description					•	
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000991_4546890_t2_174	2054	5826	150	49	}	
Description					•	
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000991_4687518_c2_519	2055	5827	585	194	179	8.0e-14
Description sp:[LN:COMK_BACSU] [AC:P40396] [GN:C TRANSCRIPTION FACTOR (CTF) (COMPETEN >pir:[LN:S43611] [AC:S43611:A69604] [GN:comK] [OR:Bacillus subtilis] [D [AC:Z99109:AL009126] [PN:competence autoregulatory control switch prior [DE:Bacillus subtilis complete genom [SP:P40396] [LE:117081] [RE:117659] [AC:S70734] [GN:comK] [OR:Bacillus s [DB:genpept-bct1] [DE:comK [Bacillus [NT:This sequence comes from Fig. 3.	CE PROTE [PN:con B:pir2] transcr to] [OR e (sect [DI:dire ubtilis]	EIN K) mpeten ygp:[0 iption :Bacil ion 6 0 ect] >0] [SR:] is, E20] [SP:Pce tran GI:e118 factor lus sub of 21): gp:[GI: Bacillu	40396] script 3044:g (CTF) tilis] from g54691 s subt mic, 3	[DB:sion fa 263337] [GN: [DB:g 999501 .7] [LN ilis E genes	wissprot] ctor (CTF) comK] [8] [LN:BSUB0006] comK] [FN:final enpept-bct1] to1209940.] [:S70734]
ORF Name AI7503000991 4688750 f3 283	NTID 2056	<u>AAID</u> 5828	<u>NT</u> Length		score	probability
	2036	3028		57		
Description						

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000991_4726010_c1_413	2057	5829	2304	767		3.7e-183
Description gp:[GI:g1196907] [LN:STMDRRC] [AC:L7 [GN:drrC] [OR:Streptomyces peucetius daunorubicin resistance protein (drr [DI:direct]] [DB:g	enpept	-bct1]	[DE:St	repton	nyces peucetius
ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	score	probability
A17503000991_4781275_f2_194	2058	5830		82		
Description NO-HIT						
ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	score	probability
AI7503000991_4812675_c1_412	2059	5831	321	106	273	3.1e-23
[DE:PROTEIN 11) (VEG11)] [SP:P54547] [PN:probable glucose-6-phosphate 1-d. [CL:glucose-6-phosphate dehydrogenas. >gp:[GI:d1013296:g1303961] [LN:BACJH subtilis] [SR:Bacillus subtilis (str. [DE:Bacillus subtilis DNA, 283 Kb re. [RE:240494] [DI:complement] >gp:[GI:[AC:Z99116:AL009126] [GN:yqjJ] [FN:u. [DE:Bacillus subtilis complete genom. [NT:similar to glucose-6-phosphate 1. [DI:direct]	ehydroge e] [OR:F 642] [AC ain:JH64 gion cor e1185654 nknown] e (secti	enase, Bacill C:D844 12(trp ntaini 1:g263 [OR:B	yqjJ] us subt 32:D823 C2 PheA ng skin 4820] [acillus of 21)	[GN:yqilis] 70] [P 1)) DN eleme LN:BSU subti : from	[jJ] [EC:1. N:YqjJ [A] [DE:nt.] [B0013] lis] [123952	1.1.49] [DB:pir2] [I] [OR:Bacillus [B:genpept-bct1] [LE:239025] [DB:genpept-bct1] [61to 2613730.]
ORF Name	NTID	AAID	NT Length	<u>AA</u> Length	score	probability
A17503000991_48587_f3_381	2060	5832	1062	353	729	4.2e-72
Description sp:[LN:GLPQ_BACSU] [AC:P37965] [GN:GLPQ] [OR:BACILLUS SUBTILIS] [EC:3.1.4.46] [DE:(GLYCEROPHOSPHODIESTER PHOSPHODIESTERASE)] [SP:P37965] [DB:swissprot] >pir:[LN:I40418] [AC:I40418:E69634:S37251] [PN:glycerophosphoryl diester phosphodiesterase glpQ] [GN:glpQ] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:d1034076:g3599635] [LN:AB006424] [AC:AB006424] [GN:ybeD] [OR:Bacillus subtilis] [SR:Bacillus subtilis (strain:168) DNA] [DB:genpept-bct1] [DE:Bacillus subtilis genomic DNA, 70 kb region between 17 and 23degree.] [LE:36004] [RE:36885] [DI:complement] >gp:[GI:g403373] [LN:BSGLPTQ] [AC:Z26522] [PN:glycerophosphoryl diester phosphodiesterase] [GN:glpQ] [FN:hydrolysis of deacylated phospholipids] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:B.subtilis glpT and glpQ genes for glycerol 3-phosphate permeaseand glycerophosphoryl diester phosphodiesterase.] [SP:P37965] [LE:1748] [RE:2629] [DI:direct] >gp:[GI:e1182165:g2632499] [LN:BSUB0002] [AC:Z99105:AL009126] [PN:glycerophosphoryl diester phosphodiesterase] [GN:glpQ] [FN:hydrolysis of deacylated phospholipids] [OR:Bacillus subtilis] [DB:genpept-bct1] [EC:3.1.4.46] [DE:Bacillus subtilis complete genome (section 2 of 21): from 194651 to415810.] [NT:alternate gene name: ybeD] [SP:P37965] [LE:38353] [RE:39234] [DI:complement]						
ORF Name	NTID	AAID	NT	<u>AA</u>	score	probability
A17503000991_4882133_c1_435	2061	5833	LengthI	Length 49		
Description NO-HIT			التتاا			

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000991_4882806_c2_524	2062	5834	522	173	399	3.9e-37
Description pir: [LN:D69838] [AC:D69838] [PN:con [OR:Bacillus subtilis] [DB:pir2] >gp [AC:Z99109:AL009126] [GN:yisU] [FN:u [DE:Bacillus subtilis complete genom [NT:similar to hypothetical proteins >gp: [GI:e1173520:g2145387] [LN:BSY09 [DB:genpept-bct1] [DE:B.subtilis 54k] [RE:23869] [DI:complement]	:[GI:e1 nknown] e (sect] [LE:1 476] [A	183088 [OR:B ion 6 65981] C:Y094	:g26334 acillus of 21) [RE:16 76] [PI	422] [I s subti : from 66643] N:YisU]	N:BSUI llis] 99950 [DI:co	B0006] [DB:genpept-bct1] 1 to1209940.] omplement] Bacillus subtilis]
ORF Name AI7503000991_4961000_f1_6 Description	NTID 2063	<u>AAID</u> 5835	<u>NT</u> Length	AA Length	score	probability
NO-HIT						
ORF Name	NTID	AAID		<u>AA</u> Length	•	probability
A17503000991_5085003_f1_118 Description	2064	5836	1455	484	1439	2.4e-147
pir: [LN:C69589] [AC:C69589] [PN:argininosuccinate lyase argH] [GN:argH] [CL:argininosuccinate lyase] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:el184193:g2635409] [LN:BSUB0015] [AC:Z99118:AL009126] [PN:argininosuccinate lyase] [GN:argH] [FN:arginine biosynthesis] [OR:Bacillus subtilis] [DB:genpept-bct1] [EC:4.3.2.1] [DE:Bacillus subtilis complete genome (section 15 of 21): from 2795131to 3013540.] [LE:215693] [RE:217078] [DI:complement] >gp:[GI:el185817:g2635428] [LN:BSUB0016] [AC:Z99119:AL009126] [PN:argininosuccinate lyase] [GN:argH] [FN:arginine biosynthesis] [OR:Bacillus subtilis] [DB:genpept-bct1] [EC:4.3.2.1] [DE:Bacillus subtilis complete genome (section 16 of 21): from 2997771to 3213410.] [LE:13053] [RE:14438] [DI:complement] >gp:[GI:g2293243] [LN:AF008220] [AC:AF008220] [PN:arginine succinate lyase] [GN:argH] [OR:Bacillus subtilis] [DB:genpept-bct2] [DE:Bacillus subtilis rrnB-dnaB genomic region.] [LE:165989] [RE:167374] [DI:direct]						
ORF Name	NTID	AAID	NT	<u>AA</u>	score	probability
Description pir:[LN:S76520] [AC:S76520] [PN:hype 6803, , PCC 6803] [SR:PCC 6803,] [DI [AC:D64002:AB001339] [PN:2-succinyl-c [SR:Synechocystis sp. (strain:PCC680: PCC6803 complete genome, 21/27, 2644* [RE:27108] [DI:complement]	B:pir2] 6-hydro: 3) DNA]	>gp:[0 xy-2,] [DB:g	tein] GI:d101 [GN:me	[OR:Syn 11017:g enD] [O	echocy 100163 R:Syne	B5] [LN:SYCSLRD] echocystis sp.] mechocystis sp.
ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probability
Description pir:[LN:A69844] [AC:A69844] [PN:hyposubtilis] [DB:pir2] >gp:[GI:e1183176] [GN:yjbI] [FN:unknown] [OR:Bacillus somplete genome (section 7 of 21): file[DI:complement]	g263351 subtilis	10] [Li s] [DB	447 tein yj N:BSUBO	148 [bI] [G 007] [pt-bct1	338 N:yjbI AC:Z99] [DE:	9110:AL009126] Bacillus subtilis

ORF Name	NTID	AAID	NT LengthL	AA ength	probability
A17503000991_5250258_c1_403	2067	5839	1839 6	1741 2	2.4e-179
Description gp:[GI:d1014255:g1651216] [LN:D88209 licheniformis] [SR:Bacillus lichenif [DE:Bacillus licheniformis DNA for F [DI:direct]	ormis (s	strain	:N22) DN	A] [DB:genpe	ept-bct1]
ORF Name AI7503000991_579441_c1_386 Description NO-HIT	NTID 2068	<u>AAID</u> 5840		AA score	probability
ORF Name AI7503000991_580307_f2_246 Description NO-HIT	NTID 2069	<u>AAID</u> 5841	<u>NT</u> LengthLe		probability
AI7503000991_6057943_c2_517	llus suk GN:yhfJ] lis comp crotein l 83] [AC:	5842 otein lotilis] [FN:: olete gligase] :Y14083	ligase home (some	omolog yhfJ] r2] >gp:[GI: [OR:Bacillu section 6 of 131] [RE:100 ypothetical tilis chromo	ell83027:g2633361] as subtilis] 5 21): from 999501 0126] [DI:direct] protein] [GN:yhfJ] osomal DNA, region
ORF Name A17503000991_6688757_f2_123 Description gp:[GI:g2267240] [LN:SEU71377] [AC:U [DB:genpept-bct1] [DE:Staphylococcus putativetranscriptional regulator At [RE:1383] [DI:direct]	epiderm	[OR:Stanidis a	aphyloco autolysia	68 823 4 ccus epiderm	

ORF Name	NTID	AAID	NT AA score probability
AI7503000991_6921877_c1_410	2072	5844	219 72 192 2.3e-14
Description sp:[LN:G6PD_BACSU] [AC:P54547] [GN:Z [DE:PROTEIN 11) (VEG11)] [SP:P54547]			
>gp:[GI:d1013296:g1303961] [LN:BACJF subtilis] [SR:Bacillus subtilis (str [DE:Bacillus subtilis DNA, 283 Kb re [RE:240494] [DI:complement] >gp:[GI: [AC:Z99116:AL009126] [GN:yqjJ] [FN:u [DE:Bacillus subtilis complete genom	se] [OR: 1642] [A rain:JH6 egion co el18565 unknown] ne (sect	Bacill C:D844 42(trp ntaini 4:g263 [OR:B	us subtilis] [EC:1.1.1.49] [DB:pir2] 32:D82370] [PN:YqjJ] [OR:Bacillus C2 PheA1)) DNA] [DB:genpept-bct1] ng skin element.] [LE:239025] 4820] [LN:BSUB0013] acillus subtilis] [DB:genpept-bct1]
ORF Name	NTID	AAID	NT AA LengthLength score probability
AI7503000991_7032752_f3_268	2073	5845	795 264 1335 2.5e-136
Description gp:[GI:g2267239] [LN:SEU71377] [AC:U [DB:genpept-bct1] [DE:Staphylococcus putativetranscriptional regulator At [DI:direct]	epider	midis	autolysin AtlE and
ORF Name	NTID	AAID	NT AA score probability
A17503000991_7081712_f1_27	2074	5846	234 77 84 0.00093
Description pir: [LN:H69835] [AC:H69835] [PN:hypsubtilis] [DB:pir2] >gp: [GI:e1183043 [GN:yhzC] [FN:unknown] [OR:Bacillus complete genome (section 6 of 21): f [DI:complement]	:g26333 subtili	77] [L s] [DB	N:BSUB0006] [AC:Z99109:AL009126] :genpept-bct1] [DE:Bacillus subtilis
ORF Name	NTID	AAID	NT AA score probability
A17503000991_7145260_f3_337 Description	2075	5847	243 80

NT AΑ ORF Name NTID LengthLength AI7503000991 7228438 c2 471 5848 3231 1076 2076 7.4e-185 Description sp:[LN:ADDA BACSU] [AC:P23478] [GN:ADDA] [OR:BACILLUS SUBTILIS] [DE:ATP-DEPENDENT NUCLEASE SUBUNIT A] [SP:P23478] [DB:swissprot] >pir:[LN:B39432] [AC:B39432:H69582] [PN:ATP-dependent deoxyribonuclease chain A addA:ATP-dependent exonuclease synthesis protein AddA] [GN:addA] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:g142440] [LN:BACADDAA] [AC:M63489] [PN:ATP-dependent nuclease] [GN:addB] [OR:Bacillus subtilis] [SR:B.subtilis DNA] [DB:genpept-bct1] [DE:Bacillus subtilis ATP-dependent nuclease (addA) and (addB), andopen reading frame 3, partial cds.] [LE:3988] [RE:7687] [DI:direct] >gp:[GI:e1183065:g2633399] [LN:BSUB0006] [AC:Z99109:AL009126] [PN:ATP-dependent deoxyribonuclease (subunit A)] [GN:addA] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 6 of 21): from 999501 to1209940.] [NT:alternate gene name: recE5] [SP:P23478] [LE:139780] [RE:143478] [DI:direct] >qp:[GI:e1364881:q2226192] [LN:BSY14081] [AC:Y14081] [PN:ATP-dependent nuclease] [GN:addA] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis chromosomal DNA, region 92 degrees: regionbetween comK and addAB.] [NT:see ref [3]; In EMBL entry M63489 this gene is] [LE:22075] [RE:25773] [DI:direct] NT AΑ LengthLength score ORF Name NTID AAID probability A17503000991 7241300 c2 502 2077 5849 135 44 Description NO-HIT AA ORF Name NTID AAID probability score LengthLength A17503000991 7308375 c3 567 2078 942 Description pir:[LN:H69668] [AC:H69668:S15231:B38447] [PN:oligopeptide transport system permease protein oppB:sporulation initiation protein spo0KB] [GN:oppB:spo0KB] [CL:oligopeptide permease protein oppB] [OR:Bacillus subtilis] [DB:pir2] >qp:[GI:e1183164:q2633498] [LN:BSUB0007] [AC:Z99110:AL009126] [PN:oligopeptide ABC transporter (permease)] [GN:oppB] [FN:required for initiation of sporulation,] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 7 of 21): from 1194391to 1411140.] [NT:alternate gene name: spo0KB] [LE:26676] [RE:27611] [DI:direct] AA ORF Name NTID AAID probability score LengthLength A17503000991 782816 cl 396 2079 1857 618 3.7e-116

Description

pir: [LN:C69975] [AC:C69975] [PN:acyltransferase homolog yrhL] [GN:yrhL] [OR:Bacillus subtilis] [DB:pir2] >gp: [GI:g1934616] [LN:BSU93874] [AC:U93874] [PN:hypothetical protein YrhL] [GN:yrhL] [OR:Bacillus subtilis] [DB:genpept-bctl] [DE:Bacillus subtilis cysteine synthase (yrhA), cystathioninegamma-lyase (yrhB), YrhC (yrhC), YrhD (yrhD), formate dehydrogenasechain A (yrhE), YrhF (yrhF), formate dehydrogenase (yrhG), YrhH(yrhH), regulatory protein (yrhI), cytochrome P450 102 (yrhJ), YrhK(yrhK), hypothetical protein YrhL (yrhL), putative anti-SigV factor(yrhM), RNA polymerase sigma factor SigV (sigV) and YrhO (yrhO)genes, complete cds, and YrhP (yrhP) gene, partial cds.] [NT:similar to Haemophilus influenzae hypothetical] [LE:13904] [RE:15808] [DI:complement] >gp: [GI:e1183944:g2635160] [LN:BSUB0014] [AC:Z99117:AL009126] [GN:yrhL] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bctl] [DE:Bacillus subtilis complete genome (section 14 of 21): from 2599451to 2812870.] [NT:similar to acyltransferase] [LE:171138] [RE:173042] [DI:direct]

ORF Name	NTID	AAID I	<u>NT</u> Length	<u>AA</u> Length	score	probability
A17503000991_813412_c1_411	2080				191	3.0e-14
Description sp: [LN:G6PD_BACSU] [AC:P54547] [GN:2] [DE:PROTEIN 11) (VEG11)] [SP:P54547] [PN:probable glucose-6-phosphate 1-6] [CL:glucose-6-phosphate dehydrogenas >gp: [GI:d1013296:g1303961] [LN:BACJF subtilis] [SR:Bacillus subtilis (str [DE:Bacillus subtilis DNA, 283 Kb re [RE:240494] [DI:complement] >gp: [GI: [AC:Z99116:AL009126] [GN:yqjJ] [FN:U] [DE:Bacillus subtilis complete genom [NT:similar to glucose-6-phosphate 1] [DI:direct]	[DB:sw lehydrog se] [OR: 1642] [A sain:JH6 sgion co e118565 unknown]	issprot enase, Bacillus C:D84433 42(trpC ntaining 4:g26348 [OR:Bac ion 13 c] >pir yqjJ] s subt 2:D823 2 PheA g skin 820] [cillus of 21)	:[LN:B6 [GN:yq] ilis] 70] [PN 1)) DNA elemen LN:BSUE subtil : from	59964] jJ] [EC:1. N:Yqj5 A] [DE nt.] 30013] Lis] 23952	[AC:B69964] 1.1.49] [DB:pir2] J] [OR:Bacillus 3:genpept-bct1] [LE:239025] [DB:genpept-bct1] 261to 2613730.]
ORF Name A17503000991_859838_f3_357 Description NO-HIT	NTID 2081			<u>AA</u> Length ¹	score	probability
ORF Name	NTID	AAID T	<u>NT</u> Length]	AA Length	score	probability
A17503000991_978436_f2_126 Description sp:[LN:PATA_BACSU] [AC:P16524] [GN:F		5854 [OR:BA	1188 ACILLU	395 S SUBTI	LLIS)	
[DE:PUTATIVE AMINOTRANSFERASE A,] [S [LN:BS16829KB] [AC:AJ222587] [PN:Pat [DB:genpept-bct1] [DE:Bacillus subti gene.] [NT:hisC homologue] [SP:P1652	A prote lis 29k	in] [GN B DNA fı	:patA] ragmen	[OR:Ba	cillu ykwC	s subtilis] gene to cse15
ORF Name	NTID	AAID L	NT ength1	<u>AA</u> Length	score	probability
A17503000991_985887_c3_570	2083	5855	435	144	570	2.9e-55
Description pir:[LN:D69843] [AC:D69843] [PN:con [CL:hypothetical protein yjbD] [OR:E >gp:[GI:el183170:g2633504] [LN:BSUB0 [OR:Bacillus subtilis] [DB:genpept-b 7 of 21): from 1194391to 1411140.] [[RE:33174] [DI:direct]	acillus 007] [Acctl] [D	subtili C:Z9911(E:Bacill	is] [Di D:AL00: lus sul	B:pir2] 9126] btilis	[GN:yj compl	bD] [FN:unknown] ete genome (section
ORF Name	NTID	AAID L	NT ength I	AA Length	core	probability
AI7503000991_9927077_£3_378	2084			48		
Description NO-HIT	<u> </u>	L	·			
ORF Name	NTID	AAID L	<u>NT</u> ength1	<u>AA</u> Length	score	probability
A17503000992_10008513_c3_117	2085	5857	2667	888	3361	0.0
Description						
<pre>pir:[LN:SYBSVS] [AC:A26738] [PN:val [CL:valinetRNA ligase] [OR:Bacillu</pre>		_		_	_	

ORF Name	NTID	AAID	NT AA score probability
AI7503000992 10720877_c1_78	2086	5858	
Description			
pir:[LN:F69723] [AC:F69723] [PN:tri [CL:trigger factor] [OR:Bacillus sub [LN:BSUB0015] [AC:Z99118:AL009126] [tilis] PN:trig ctl] [D [NT:alt] >gp:[tig] [O 9bp.] [[DB:pi ger fa E:Baci ernate GI:e11 R:Baci NT:hom	r2] >gp:[GI:e1184072:g2635288] ctor (prolyl isomerase)] [GN:tig] llus subtilis complete genome (section gene name: yzzH] [SP:P80698] 65373:g1770074] [LN:BSZ75208] llus subtilis] [DB:genpept-bct1] ology to trigger factor of
ORF Name	NTID	AAID	NT AA LengthLength score probability
AI7503000992_11954500_f3_71	2087	5859	174 57
Description NO-HIT			
ORF Name	NTID	AAID	NT AA score probability
A17503000992_1211562_c2_94	2088	5860	1269 422 1518 1.0e-155
Description gp:[GI:e221213:g1296452] [LN:BSCLPXG [OR:Bacillus subtilis] [DB:genpept-b protease] [SP:P50866] [LE:168] [RE:1	ct1] [D	E:B.su	btilis clpX gene.] [NT:ATP-dependent
ORF Name	NTID	AAID	NT AA score probability
AI7503000992_14725300_c3_108	2089	5861	123 40
Description NO-HIT			
ORF Name	NTID	AAID	NT <u>AA</u> LengthLength
AI7503000992_15820252_c1_81	2090	5862	141 46
Description		-	
NO-HIT			
ORF Name	NTID	AAID	NT AA LengthLength
A17503000992_16835900_c2_104	2091	5863	147 48 74 0.011
Description			
gp:[GI:g1123053] [LN:CELF59A6] [AC:U	41994]	[GN:F5	9A6.2] [OR:Caenorhabditis elegans]

[SR:Caenorhabditis elegans strain=Bristol N2] [DB:genpept-inv1] [DE:Caenorhabditis

elegans cosmid F59A6.] [LE:19719:20072] [RE:19904:20242] [DI:directJoin]

		NT AA
ORF Name	<u>NTID</u>	AAID LengthLength score probability
A17503000992_188388_c3_122	2092	5864 579 192 220 3.6e-18
Description		
sp:[LN:MREC_BACSU] [AC:Q01466] [GN:M		
		[DB:swissprot] >pir:[LN:JC4595] [AC:JC4595
<pre>[PN:cell shape determinant MreC:pr [GN:mreC] [OR:Bacillus stearothermone</pre>		
		eterminant mreC] [GN:mreC] [OR:Bacillus
		CMREMIN] [AC:M95582] [GN:mreC] [FN:cell
		SR:Bacillus subtilis (strain W168) DNA]
[DB:genpept-bct1] [DE:Bacillus subti		
completecds, (mred) gene complete co cds.] [NT:putative] [LE:1370] [RE:22		C) gene complete cds, (minD)gene complete
cus.; [MI.putative] [HE.1370] [RE.22	.42] [DI	.uirectj
ORF Name	NTID	AAID NT AA score probability
AI7503000992_19707767_f1_4	2093	5865 324 107
Description		
NO-HIT		
		NT AA
ORF Name	NTID	AAID LengthLength score probability
AI7503000992_20348160_f1_1	2094	5866 222 73 71 0.022
Description		
		17055:U32570] [GN:A130R] [OR:Paramecium
] [DE:Paramecium bursaria Chlorella virus 1,
complete genome.] [LE:69061] [RE:693	148] [DI	:direct)
ORF Name	NTID	AAID NT AA score probability
A17503000992 21500253 c3 109	BARE	LengthLength
	2095	5867 177 58
Description NO-HIT		
NO-HII		
ORF Name	NTID	AAID NT AA score probability
A17503000992_21563137_c2_102		<u> </u>
	2096	5868 135 44
Description NO-HIT		
NO-III	****	
ORF Name	NTID	AAID Longth ongth score probability
A17503000992 23444187 f3 62		Lengthength
Description	2097	5869 162 53
Deport De Toll		

ORF Name	NTID	AAID	NT AA score probability
A17503000992_23522567_c2_91	2098	5870	381 126 450 1.5e-42
Description			
subtilis] [DB:pir2] >gp:[GI:e1184134 [PN:ribosomal protein L20] [GN:rplT] [DE:Bacillus subtilis complete genom [SP:P55873] [LE:156202] [RE:156561]	ot] >pi: herichia :g26353! [OR:Bad e (sect: [DI:comp mal proto omic sec	r:[LN: a coli 50] [L cillus ion 15 plemen tein L quence	EF69696] [AC:F69696] [PN:ribosomal iribosomal protein L20] [OR:Bacillus LN:BSUB0015] [AC:Z99118:AL009126] s subtilis] [DB:genpept-bct1] s of 21): from 2795131to 3013540.] at] >gp:[GI:e1165301:g1770009] L20] [GN:rplT] [OR:Bacillus subtilis] s 89009bp.] [NT:homology to rplT of
ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000992_23531628_£2_28	2099	5871	144 47
Description NO-HIT		414/1	
ORF Name	NTID	AAID	NT AA score probability
A17503000992_23697141_c2_97	2100	5872	930 309 1327 1.8e-135
<pre>gp:[GI:g2589181] [LN:SAU89396] [AC:U [OR:Staphylococcus aureus] [DB:genpe] hemCDBL gene cluster: porphobilinoge: (hemD),d-aminolevulinic acid dehydra: complete cds.] [LE:219] [RE:1145] [D</pre>	pt-bct2] ndeamina tase (he] [EC: ase (h emB) a	4.3.1.8] [DE:Staphylococcus aureus
ORF Name	NTID	AAID	NT AA score probability
A17503000992_24017052_c2_103	2101	5873	150 49
Description			
NO-HIT			
ORF Name	NTID	AAID	NT AA score probability
AI7503000992_24240877_c3_119	2102	5874	522 173
Description			
NO-HIT			
ORF Name	NTID	AAID	NT AA score probability
A17503000992_24625161_c1_87	2103	5875	141 46 160 8.2e-12
Description			
<pre>sp:[LN:RADC_STAAU] [AC:P31337] [GN:RADC HOMOLOG (25 KD PROTEIN) >gp:[GI:g551992] [LN:STATNIS5] [AC:KG (strain RN450) DNA] [DB:genpept-bct1] insertion site.] [NT:25 kD protein ()</pre>	(FRAGMI 02985]] [DE:S	ENT)] [OR:St .aureu	[SP:P31337] [DB:swissprot] caphylococcus aureus] [SR:S.aureus

ORF Name	NTID	AAID	NT LengthL	ength scor	e probability	
AI7503000992_24897312_c3_112	2104	5876		51 255	7.1e-22	
Description sp:[LN:HEM2_STAAU] [AC:P50915] [GN:HEM2_STAAU] [AC:P50915] [GN:HEM2_STAAU] [AC:P50915] [GN:HEM2_SYNTHASE) (ALAD) (ALADH)] [SP:P500000000000000000000000000000000000	0915] [olevuli pt-bct2 ndeamin tase (h	DB:swisnic act nic act] [EC:4 ase (he emB) ar	ssprot] id dehyd 1.2.1.24 emC), ur	>gp:[GI:gi lratase] [G l] [DE:Stap coporphyrin	2589183] EN:hemB] phylococcus aur nogen III synth	ase
ORF Name	NTID	AAID	<u>NT</u> LengthL	AA ength	e probability	
AI7503000992_25582912_£2_23	2105	5877		10		
Description NO-HIT						
ORF Name 417503000992 25596000 c3 111	NTID	<u>AAID</u> 5878		AA ength	e probability	
Description sp:[LN:HEMX_BACSU] [AC:P16645] [GN:HI [SP:P16645] [DB:swissprot] >pir:[LN:I negative effector hemX] [GN:hemX] [GI [LN:BACHEMAXC] [AC:M57676:M32130] [GI [DB:genpept-bct1] [DE:Bacillus subtil product] [LE:1607] [RE:2437] [DI:dire [AC:Z99118:AL009126] [PN:membrane-box concentration of HemA] [OR:Bacillus subtil complete genome (section 15 of 21): [RE:81728] [DI:complement] >gp:[GI:ei [PN:membrane-bound protein] [GN:hemX] [DE:B.subtilis genomic sequence 89008 heme] [SP:P16645] [LE:88179] [RE:8908	EMX] [OB35252] OR:Baci N:hemX] lis hem ect] >g und pro subtili from 27 1165382] [OR:B	R:BACII [AC:B3 llus su [OR:Ba AXCDBL p:[GI:e tein] s] [DB: 95131tc :g1770c acillus NT:memb	LUS SUE 35252:C6 acillus gene cl 21184065 [GN:hemX:genpept 301354 082] [LN s subtil	TILIS] [DI [DB:pir2] subtilis] [uster.] [I [:g2635281] [] [FN:nega [:-bct1] [DI [:0.] [SP:PI [:BSZ75208] [is] [DB:ge	E:HEMX PROTEIN] N:hemA concentr >gp:[GI:g1430 [SR:B.subtilis NT:unidentified [LN:BSUB0015] Ative effector E:Bacillus subt [A6645] [LE:8089 [AC:Z75208] Enpept-bct1]	36] DNA] gene of the ilis
ORF Name	NTID			AA ength		
A17503000992_2866255_f3_65	2107	5879	264	74	0.045	
Description gp:[GI:e1250026:g2842472] [LN:SPBC20] [GN:SPBC20F10.10] [OR:Schizosaccharor [DE:S.pombe chromosome II cosmid c20] SIMILARITY:Arabidopsis] [LE:20474] [I	myces p F10.] [ombe] (NT:SPBC	SR:fiss 20F10.1	ion yeast] 0, len:243	[DB:genpept-p]	ln1]
ORF Name	NTID	AAID	<u>NT</u> LengthL	AA ength	probability	
A17503000992_31510_f3_55	2108	5880	627 2	108 435	6.0e-41	
Description sp:[LN:3MGA_HAEIN] [AC:P44321] [GN:TR [EC:3.2.2.20] [DE:GLYCOSYLASE) (TAG)] [AC:G64084] [PN:3-methyladenine DNA glycosylase I] [OR:Haemophilus influe [LN:U32748] [AC:U32748:L42023] [PN:DR [GN:HI0654] [OR:Haemophilus influenza Rd section 63 of 163 of the complete GB:X03845 PID:147920] [LE:6773] [RE:	[SP:Postenzae] NA-3-me ae Rd] genome	44321] ylase, [EC:3.2 thylade [DB:gen	[DB:swi I] [CL: 2.2] [nine gl pept-bc similar	ssprot] >p 3-methylad DB:pir2] > ycosidase t2] [DE:Ha	oir:[LN:G64084] denine DNA gp:[GI:g157365; I (tagI)] demophilus influ	

			1777	7.70	
ORF Name	NTID	AAID	NT LengthL	AA ength score	probability
AI7503000992_33395050_c2_95	2109	5881		05 577	5.3e-56
Description	L				<u> </u>
sp:[LN:YSXC_BACSU] [AC:P38424] [GN:Y [DB:swissprot] >pir:[LN:I40422] [AC:protein ysxC] [GN:ysxC] [CL:Bacillu ysxC:translation elongation factor T >gp:[GI:g496558] [LN:BSLONLA] [AC:X7 [DE:B.subtilis lon gene for protease [DI:direct] >gp:[GI:e1184068:g263528 [FN:unknown] [OR:Bacillus subtilis] genome (section 15 of 21): from 2795 proteins] [SP:P38424] [LE:83852] [RE [LN:BSZ75208] [AC:Z75208] [PN:hypoth [DB:genpept-bct1] [DE:B.subtilis gen putative] [SP:P38424] [LE:85468] [RE	I40422:0 Is subtiling the subtiling of t	C69987 lis co ogy] [OR:Bac NT:orf BSUB00 pept-b 013540 [DI:c protei quence	:S45102 nserved OR:Bacil illus su X] [SP:P 15] [AC: ct1] [DE .] [NT:s omplemen n] [GN:y 89009bp] [PN:conshypothetic lus subtil stilis] [D 38424] [LE Z99118:AL0:Bacillus imilar to t] >gp:[GI sxC] [OR:B	erved hypothetical al protein is] [DB:pir2] B:genpept-bct1] :2669] [RE:3256] 09126] [GN:ysxC] subtilis complete hypothetical :e1165379:g1770079] acillus subtilis]
ORF Name	NTID	AAID	NT LengthLe	AA ength	probability
AI7503000992_33517_c1_82	2110	5882	1077 3	58 258	2.8e-21
Description					
pir:[LN:E69834] [AC:E69834] [PN:hyp subtilis] [DB:pir2] >gp:[GI:e1183059 [GN:yhjN] [FN:unknown] [OR:Bacillus complete genome (section 6 of 21): f [DI:complement] >gp:[GI:e324987:g222 protein] [GN:yhjN] [OR:Bacillus subt chromosomal DNA, region 92 degrees: Similarity to a hypothetical protein	:g263333 subtilis rom 9995 6186] [I ilis] [I regionbe	93] [Lis] [DB 501 to LN:BSY DB:gen etween	N:BSUB00 :genpept 1209940. 14081] [pept-bct comK and	06] [AC:Z9 -bct1] [DE] [LE:1308 AC:Y14081] 1] [DE:Bac d addAB.]	9109:AL009126] :Bacillus subtilis 91] [RE:132045] [PN:hypothetical illus subtilis [NT:TTG start;
ORF Name	NTID	AAID	NT LengthLe	AA ength	probability
A17503000992_34642092_c1_85	2111	5883	1284 4	27 907	5.7e-91
Description					
<pre>sp:[LN:FOLC_BACSU] [AC:Q05865] [GN:F [DE:SYNTHETASE) (FPGS)] [SP:Q05865] [AC:B40646:B33490:D69626] [PN:folyl [OR:Bacillus subtilis] [DB:pir2] >gp [PN:folyl-polyglutamate synthetase] subtilis (sub_strain PY79, strain W1</pre>	[DB:swis -polyglu :[GI:g14 [GN:fold	ssprot ltamate 12936] C] [OR	>pir:[] = synthet [LN:BAC] :Bacillus	LN:B40646] case folc] FOLC] [AC: s subtilis	[GN:folC] L04520] [SR:Bacillus

valyl tRNA synthetase (valS) gene, 3' end cds;folyl-polyglutamate synthetase (folC) gene, complete cds; comCgene, 5' end cds.] [LE:231] [RE:1523] [DI:direct] >gp:[GI:e1184057:g2635273] [LN:BSUB0015] [AC:Z99118:AL009126] [PN:folyl-polyglutamate synthetase] [GN:folC] [FN:folate biosynthesis] [OR:Bacillus subtilis] [DB:genpept-bct1] [EC:6.3.2.17] [DE:Bacillus subtilis complete genome (section 15 of 21): from 2795131to 3013540.] [SP:Q05865] [LE:69443] [RE:70735] [DI:complement]

NT AAORF Name NTID AAID score probability LengthLength A17503000992_3914012_c2_98 2112 5884 690

Description

gp:[GI:g2589182] [LN:SAU89396] [AC:U89396] [PN:uroporphyrinogen III synthase] [GN:hemD] [OR:Staphylococcus aureus] [DB:genpept-bct2] [EC:4.2.1.75] [DE:Staphylococcus aureus hemCDBL gene cluster: porphobilinogendeaminase (hemC), uroporphyrinogen III synthase (hemD), d-aminolevulinic acid dehydratase (hemB) and GSA-1-aminotransferase(hemL) genes, complete cds.] [LE:1167] [RE:1835] [DI:direct]

ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000992_4080443_c3_113	2113	5885	1296 431 2041 3.9e-211
Description			
	TRANSFER :U89396] ept-bct2 endeamina atase (he	ASE) ([PN:G] [EC: ase (h emB) a	GSA-AT)] [SP:034092] [DB:swissprot] SA-1-aminotransferase] [GN:hemL]
ORF Name	NTID	AAID	NT AA score probability
AI7503000992_439183_f1_14	2114	5886	297 98 208 6.8e-17
Description	-	<u> </u>	
<pre>gp:[GI:e1165375:g1770076] [LN:BSZ752 [OR:Bacillus subtilis] [DB:genpept-} [NT:unknown function; putative] [LE</pre>	bct1] [D]	E:B.su	
ORF Name	NTID	AAID	NT AA score probability
AI7503000992_4729837_c2_92	2115	5887	624 207 416 6.1e-39
Description			
KD PROTEIN IN NRDF-CWLC INTERGENIC F [AC:D69883] [PN:hypothetical protein >gp:[GI:e218063:g1154634] [LN:BSNRDM [OR:Bacillus subtilis] [DB:genpept-hand genes.] [NT:no similarities, cannotation [DI:direct] >gp:[GI:e1183399:g263412] [FN:unknown] [OR:Bacillus subtilis]	REGION] in ymaB] yMA] [AC oct1] [DI be inact 24] [LN:I [DB:genp	[SP:P5 [GN:y: :Z6850 E:B.su tivate BSUB00 pept-b	btilis cwlC, nrdE, nrdF, ymaA and ymaB d] [SP:P50619] [LE:4450] [RE:5070] 10] [AC:Z99113:AL009126] [GN:ymaB]
ORF Name	NTID	AAID	NT AA LengthLength score probability
AI7503000992_4891876 f3 45	2116	5888	141 46

Description NO-HIT

ORF Name score NTID probability A17503000992 500052_c3_118 5889 669 1.8e-41 2117 Description sp:[LN:RADC BACSU] [AC:Q02170] [GN:YSXA] [OR:BACILLUS SUBTILIS] [DE:DNA REPAIR PROTEIN RADC HOMOLOG (ORFB)] [SP:Q02170] [DB:swissprot] >pir:[LN:B45239] [AC:B45239:B45240:I39881:A69987] [PN:DNA repair protein homolog ysxA] [GN:ysxA] [CL:DNA repair protein radc] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:g142854] [LN:BACDIVREG] [AC:M96343] [GN:orfB] [OR:Bacillus subtilis] [SR:Bacillus subtilis (sub_strain PY79, strain W168) DNA] [DB:genpept-bct1] [DE:Bacillus subtilis orfA, orfB, mreB, mreC, mreD, minC, and minDgenes, complete coding regions.] [NT:homologous to E. coli radC gene product and to] [LE:762] [RE:1457] [DI:direct] >gp:[GI:g143162] [LN:BACMAFMREB] [AC:L08793] [OR:Bacillus subtilis] [SR:Bacillus subtilis (strain W168) DNA] [DB:genpept-bct1] [DE:Bacillus subtilis maf gene, complete cds; orfB, complete cds; mreBgene, 5' end.] [NT:putative] [LE:1557] [RE:2252] [DI:direct] >qp: [GI:e1184053:q2635269] [LN:BSUB0015] [AC:Z99118:AL009126] [GN:ysxA] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 15 of 21): from 2795131to 3013540.] [NT:similar to DNA repair protein] [SP:Q02170] [LE:65971] [RE:66666] [DI:complement] NT AΑ ORF Name NTID AAID <u>sco</u>re probability LengthLength A17503000992 5268775 c3 105 2118 5890 960 319 8.5e-07 Description pir:[LN:H69986] [AC:H69986] [PN:hypothetical protein ysoA] [GN:ysoA] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1184073:g2635289] [LN:BSUB0015] [AC:Z99118:AL009126] [GN:ysoA] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 15 of 21): from 2795131to 3013540.] [LE:91788] [RE:92720] [DI:complement] >gp:[GI:e1165372:g1770073] [LN:BSZ75208] [AC:Z75208] [PN:hypothetical protein] [GN:ysoA] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:B.subtilis genomic sequence 89009bp.] [NT:unknown function; putative] [LE:77187] [RE:78119] [DI:direct] NT AA ORF Name NTID AAID score probability LengthLength AI7503000992 5860827 c3 110 2119 5891 1347 448 1172 4.8e-119 Description

sp:[LN:HEM1 BACSU] [AC:P16618] [GN:HEMA] [OR:BACILLUS SUBTILIS] [EC:1.2.1.-] [DE:GLUTAMYL-TRNA REDUCTASE, (GLUTR)] [SP:P16618] [DB:swissprot] >pir:[LN:A35252] [AC:A35252:C69639] [PN:glutamyl-tRNA reductase, hemA:hemA protein] [GN:hemA] [CL:glutamyl-tRNA reductase] [OR:Bacillus subtilis] [EC:1.2.1.-] [DB:pir2] >gp:[GI:g143035] [LN:BACHEMAXC] [AC:M57676:M32130] [PN:NAD(P)H:qlutamyl-transfer RNA reductase] [GN:hemA] [OR:Bacillus subtilis] [SR:B.subtilis DNA] [DB:genpept-bct1] [DE:Bacillus subtilis hemAXCDBL gene cluster.] [NT:The product of this hemA gene is not an] [LE:232] [RE:1599] [DI:direct] >qp:[GI:e1184066:q2635282] [LN:BSUB0015] [AC:Z99118:AL009126] [PN:glutamyl-tRNA reductase] [GN:hemA] [FN:porphyrin biosynthesis] [OR:Bacillus subtilis] [DB:genpept-bct1] [EC:1.2.1.-] [DE:Bacillus subtilis complete genome (section 15 of 21): from 2795131to 3013540.] [SP:P16618] [LE:81736] [RE:83103] [DI:complement] >gp:[GI:e1165381:g1770081] [LN:BSZ75208] [AC:Z75208] [PN:NAD(P)H:glutamyl-transfer RNA reductase] [GN:hemA] [OR:Bacillus subtilis] [DB:genpept-bct1] [EC:2.3.1.37] [DE:B.subtilis genomic sequence 89009bp.] [NT:NAD(P)H:glutamyl-t-RNA reductase of the C-5 pathway] [SP:P16618] [LE:86804] [RE:88171] [DI:direct]

ORF Name	NTID	AAID LengthLength score probability
A17503000992_6845382_£2_26	2120	5892 219 72 0.034
Description gp:[GI:e1358367:g3979946] [LN:CEY18D:	 10A] [A0	C:AL034393] [GN:Y18D10A.14]
[OR:Caenorhabditis elegans] [DB:genpe Y18D10A, complete sequence.] [LE:1138 [DI:directJoin]	ept-inv	l] [DE:Caenorhabditis elegans cosmid
ORF Name	NTID	AAID NT AA score probability
A17503000992_7050319_c2_90	2121	LengthLength 5893 237 78 226 8.4e-19
Description		
sp:[LN:RL35_BACSU] [AC:P55874] [GN:RIPROTEIN L35] [SP:P55874] [DB:swisspro		R:BACILLUS SUBTILIS] [DE:50S RIBOSOMAL
ORF Name	NTID	AAID LengthLength score probability
A17503000992_7283437_c1_89	2122	5894 162 54
Description NO-HIT		
NO-IIII		
ORF Name	NTID	AAID LengthLength score probability
A17503000992_806510_c1_86	2123	5895 714 237 189 7.0e-15
Description		
COMPETENCE PROTEIN COMC)] [SP:P15378] [AC:A33490:E40646:F40646:B69602] [Protein competence protein competence protein competence protein competence protein competence protein comc) [Protein comc) gene, competence protein comc) gene, completeds.] [In:BSUB06 [GN:comc] [FN:required for the process [DB:genpept-bct1] [DE:Bacillus subtilication competence] [DE:P15378] [LE:60]	DB:sv N:type D in ComC] [DB:pin ein] [GN ept-bct1 LE:819] D15] [AC ssing ar Lis comp	[GN:comC] [CL:type IV prepilin peptidase] [GN:comC] [CL:type IV prepilin peptidase] [C] >gp:[GI:g142704] [LN:BACCOMC] [ComC] [OR:Bacillus subtilis] [SR:Bacillus [C] [DE:Bacillus subtilis late competence [RE:1565] [DI:direct] [C:Z99118:AL009126] [PN:DNA-binding protein] [C] dtranslocation] [OR:Bacillus subtilis] [C] plete genome (section 15 of 21): from [RE:69303] [DI:complement]
ORF Name AI7503000992_820253_f1_15	NTID	Length Length score probability
Description	2124	5896 297 98 206 1.1e-16
gp:[GI:e1165375:g1770076] [LN:BSZ7520	tl] [DE	Z75208] [PN:hypothetical protein] [GN:ysoC] :B.subtilis genomic sequence 89009bp.] [RE:81206] [DI:complement]
ORF Name	NTID	AAID NT AA score probability
A17503000992_978562_c1_79	2125	5897 861 286 1261 1.8e-128
Description		
[DE:SYNTHASE) (ALAD) (ALADH)] [SP:P50 [LN:SAU89396] [AC:U89396] [PN:d-amino [OR:Staphylococcus aureus] [DB:genper hemCDBL gene cluster: porphobilinogen	915] [Dolevuling ot-bct2] adeamina (he	ic acid dehydratase] [GN:hemB] [EC:4.2.1.24] [DE:Staphylococcus aureus se (hemC), uroporphyrinogen III synthase mB) and GSA-1-aminotransferase(hemL) genes,

ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000993_10266875_c2_761 Description	2126	5898	156 51
NO-HIT			
ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000993_10463_c2_803	2127	5899	504 167 315 3.1e-28
] [OR:Bacillus subtilis] [DB:pir2] > [OR:Bacillus subtilis] [DB:genpept-b UDP-N-acetylmuramate-alanine ligase [RE:1642] [DI:direct] >gp:[GI:e11858	EGION (G [PN:gengp:[GI:got1] [Digene, passible of the passible	ORF1)] neral g55601 E:Baci artial 5462] s] [DB 97771t 48897] O] [PN	[SP:P40779] [DB:swissprot] stress protein homolog ytxG] [GN:ytxG 5] [LN:BACUNAM] [AC:L31845] llus subtilis cds, and 3 ORF's.] [NT:ORF1] [LE:1211] [LN:BSUB0016] [AC:Z99119:AL009126] :genpept-bct1] [DE:Bacillus subtilis o 3213410.] [NT:alternate gene name: [RE:49328] [DI:complement] :YtxG] [GN:ytxG] [OR:Bacillus
ORF Name A17503000993_10548383_f1_3 Description NO-HIT	NTID 2128	<u>AAID</u> 5900	NT AA score probability LengthLength 237 78
ORF Name A17503000993_10564375_t3_510 Description NO-HIT	NTID 2129	<u>AAID</u> 5901	NT AA score probability LengthLength 123 40
ORF Name	NTID	AAID	NT AA score probability
A17503000993_1056693_c2_793 Description	2130	5902	1101 366 774 7.1e-77
	pha chai 016] [AC ct1] [DI [NT:simi 2293210] :genpept	in] [0] C:Z991 E:Baci ilar to [LN::	R:Bacillus subtilis] [DB:pir2] 19:AL009126] [GN:ytoP] [FN:unknown] 1lus subtilis complete genome (section endo-1,4-beta-glucanase] [LE:56559] AF008220] [AC:AF008220] [PN:YtoP] 1 [DE:Bacillus subtilis rrnB-dnaB
ORF Name	NTID	AAID	NT AA LengthLength score probability
AI7503000993_10588877_±1_52	2131	5903	138 45
Description NO-HIT			

ORF Name	NTID	AAID NT AA score probability
A17503000993_10667002_t1_111	2132	5904 [165]54
Description		
NO-HIT		
	and the second	NT AA
ORF Name	NTID	AAID LengthLength score probability
A17503000993_109430_f3_485	2133	5905 144 47
Description		
NO-HIT		
ORF Name	NTID	AAID NT AA score probability
MT7502000902 10069427 #2 448	1313 <i>X</i> 1	<u> </u>
A17503000993_10969427_f3_448	2134	5906 156 51
Description NO-HIT		
NO-HII		
ORF Name	NTID	AAID NT AA score probability
AI7503000993_112525_f3_546	2135	5907 141 46
Description		
NO-HIT		
	energy to the second of the second of	NTT A A
ORF Name	NTID	AAID LengthLength score probability
AI7503000993_116337_f2_251	2136	5908 627 208 199 6.1e-16
Description		
gp:[GI:g3043872] [LN:LLU95837] [AC:U		
		DE:Lactococcus lactis transmembrane protein
[RE:588] [DI:complement]	iomorog;	identified as a fusion to a signal] [LE:<1]
ORF Name	NTID	AAID NT AA score probability
A17503000993_11881325_f3_568	2137	5909 147 48
Description	L	
NO-HIT		
		NT AA
ORF Name	NTID	AAID LengthLength score probability
AI7503000993_1203827_c3_893	2138	5910 639 212 475 3.4e-45
Description		
sp:[LN:RISA_ACTPL] [AC:P50854] [GN:R	IBE:RIB	B] [OR:ACTINOBACILLUS PLEUROPNEUMONIAE]
		1.9] [DE:RIBOFLAVIN SYNTHASE ALPHA CHAIN,]
<pre>[SP:P50854] [DB:swissprot] >gp:[GI:g synthase alpha subunit] [GN:ribB] [O</pre>		[LN:APU27202] [AC:U27202] [PN:riboflavin
[DB:genpept-bct1] [DE:Actinobacillus		
		, riboflavin synthase alphasubunit (ribB),
bifunctional GTP cyclohydraseII/3,4-	dihydro	xy-2-butanone-4-phosphate synthase (ribA),
andriboflavin synthase beta subunit similar to Bacillus subtilis] [LE:16		genes, complete cds.] [NT:lumazine synthase;

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	score	probabil	ity
AI7503000993_1209417_c3_959	2139	5911	1704	567	797	2.6e-79	
Description			لـــاك				
sp:[LN:PHOR_BACSU] [AC:P23545] [GN:FIDE:ALKALINE PHOSPHATASE SYNTHESIS SOPIT: [LN:A27650] [AC:A27650:G69676] phoR] [GN:phoR] [OR:Bacillus subtiliants [AC:M23549] [PN:alkaline phosphatase subtilis] [SR:Bacillus subtilis DNA] phosphatase regulatory protein (phore [RE:1824] [DI:direct] >gp:[GI:e11841] [PN:two-component sensor histidine pregulation] [OR:Bacillus subtilis] [Senome (section 15 of 21): from 2795 [DI:complement] >gp:[GI:g2293271] [Improtein kinase] [GN:phoR] [OR:Bacillus subtilis] subtilis rrnB-dnaB genomic region.]	EENSOR PR [PN:pho Lis] [DB: e regulat [DB:gene, 3" 59:g2635 cinase] [DB:genpe 6131to 30 LN:AF0082	ROTEIN Desphate pir2] tory pr pept-1 end a 5375] [GN:pha pt-bcs 013540 220] [A	PHOR,] e respo ygp:[G rotein] oct1] [and pho [LN:BSU oR] [FN t1] [DE c] [SP: AC:AF00 [DB:gen	[SP:Ponse reconstruction [SP:Ponse reconstruction [SP:Ponser]] [SP:Ponser] [SP	23545] gulato 331] [hoR] [illus , comp [AC:2 ved in lus su] [LE: [PN:si ct2] [[DB:swiss or histiding In:BACPHORE CR:Bacillus subtilis and phosphate subtilis con 180011] [Fignal trans DE:Bacillus or history or h	ne kinase RP] us alkaline [] [LE:85] 09126] e mplete RE:181750] sduction
ORF Name	NTID	AAID	$\underline{\mathtt{NT}}$	<u>AA</u>	score	probabil	ity
			Length		BCOIC	probabii	<u> </u>
A17503000993_125880_f2_361	2140	5912	309	102			
Description							
NO-HIT				·		· ·	
ORF Name	NTID		<u>NT</u> Length	<u>AA</u> Length	score	probabil	ity
AI7503000993_1284381_c1_652	2141						
	2141	5913	741	246	297	2.5e-26	
Description sp:[LN:TAL_METJA] [AC:Q58370] [GN:TAL_EC:2.2.1] [DE:TRANSALDOLASE-LIKE >pir:[LN:H64419] [AC:H64419] [PN:tr phosphoprotein orfU] [OR:Methanococc [MP:REV892437-891784] >gp:[GI:g1591 [PN:transaldolase] [GN:MJ0960] [OR:M [DE:Methanococcus jannaschii section SP:P19669 PID:853766 GB:AL009126 per	AL:MJ0960 PROTEIN, ransaldol rus janna 624] [LN lethanoco	[OR] [SP .ase,] .schii] J:U6753 occus [METHAN Q58370 [CL:Ba [EC:2 39] [AC jannasc	OCOCCUS Cillus .2.1.2 :U6753 hii] [Implete	S JANN swissp subti [DB: 9:L771 DB:gen genom	ASCHII] rot] lis 23K pir2] 17] pept-bct2] e.] [NT:si	
Description sp:[LN:TAL_METJA] [AC:Q58370] [GN:TAL [EC:2.2.1] [DE:TRANSALDOLASE-LIKE >pir:[LN:H64419] [AC:H64419] [PN:tr phosphoprotein orfU] [OR:Methanococc [MP:REV892437-891784] >gp:[GI:g1591 [PN:transaldolase] [GN:MJ0960] [OR:M [DE:Methanococcus jannaschii section SP:P19669 PID:853766 GB:AL009126 per	AL:MJ0960 PROTEIN, ransaldol rus janna 624] [LN lethanoco 81 of 1 rcent] [I	[OR] [OR] [SP] [ASE,] [SCHII]	METHAN :Q58370 [CL:Ba [EC:2 39] [AC jannasc the co D] [RE:	OCOCCUS Collus Collu	S JANN swissp subti] [DB: 9:L771 DB:gen genom [DI:co	ASCHII] rot] lis 23K pir2] 17] pept-bct2] e.] [NT:si	milar to
Description sp:[LN:TAL_METJA] [AC:Q58370] [GN:TAL [EC:2.2.1] [DE:TRANSALDOLASE-LIKE >pir:[LN:H64419] [AC:H64419] [PN:tr phosphoprotein orfU] [OR:Methanococc [MP:REV892437-891784] >gp:[GI:g1591 [PN:transaldolase] [GN:MJ0960] [OR:M [DE:Methanococcus jannaschii section SP:P19669 PID:853766 GB:AL009126 per ORF Name AT7503000993_12894378_f2_347	AL:MJ0960 PROTEIN, cansaldol cus janna .624] [LN lethanoco .81 of 1	[OR] [OR] [SP] [SP] [SP] [SP] [SP] [SP] [SP] [SP	METHAN :Q58370 [CL:Ba [EC:2 39] [AC jannasc the co D] [RE:	OCOCCUS] [DB:s cillus .2.1.2]:U67539 hii] [I mplete 8643]	S JANN swissp subti] [DB: 9:L771 DB:gen genom [DI:co	ASCHII] rot] lis 23K pir2] 17] pept-bct2] e.] [NT:si mplement]	milar to
Description sp:[LN:TAL_METJA] [AC:Q58370] [GN:TAL [EC:2.2.1] [DE:TRANSALDOLASE-LIKE >pir:[LN:H64419] [AC:H64419] [PN:tr phosphoprotein orfU] [OR:Methanococc [MP:REV892437-891784] >gp:[GI:g1591 [PN:transaldolase] [GN:MJ0960] [OR:M [DE:Methanococcus jannaschii section SP:P19669 PID:853766 GB:AL009126 per ORF Name A17503000993_12894378_f2_347 Description	AL:MJ0960 PROTEIN, ransaldol rus janna 624] [LN lethanoco 81 of 1 rcent] [I	[OR] [OR] [SP] [ASE,] [SCHII]	METHAN :Q58370 [CL:Ba [EC:2 39] [AC jannasc the co D] [RE:	OCOCCUS Collus Collu	S JANN swissp subti] [DB: 9:L771 DB:gen genom [DI:co	ASCHII] rot] lis 23K pir2] 17] pept-bct2] e.] [NT:si mplement]	milar to
Description sp:[LN:TAL_METJA] [AC:Q58370] [GN:TAL [EC:2.2.1] [DE:TRANSALDOLASE-LIKE >pir:[LN:H64419] [AC:H64419] [PN:tr phosphoprotein orfU] [OR:Methanococc [MP:REV892437-891784] >gp:[GI:g1591 [PN:transaldolase] [GN:MJ0960] [OR:M [DE:Methanococcus jannaschii section SP:P19669 PID:853766 GB:AL009126 per ORF Name AT7503000993_12894378_f2_347	AL:MJ0960 PROTEIN, ransaldol rus janna 624] [LN lethanoco 81 of 1 rcent] [I	[OR] [OR] [SP] [ASE,] [SCHII]	METHAN :Q58370 [CL:Ba [EC:2 39] [AC jannasc the co D] [RE:	OCOCCUS Collus Collu	S JANN swissp subti] [DB: 9:L771 DB:gen genom [DI:co	ASCHII] rot] lis 23K pir2] 17] pept-bct2] e.] [NT:si mplement]	milar to
Description sp:[LN:TAL_METJA] [AC:Q58370] [GN:TAL [EC:2.2.1] [DE:TRANSALDOLASE-LIKE >pir:[LN:H64419] [AC:H64419] [PN:tr phosphoprotein orfU] [OR:Methanococc [MP:REV892437-891784] >gp:[GI:g1591 [PN:transaldolase] [GN:MJ0960] [OR:M [DE:Methanococcus jannaschii section SP:P19669 PID:853766 GB:AL009126 per ORF Name A17503000993_12894378_f2_347 Description	AL:MJ0960 PROTEIN, ransaldol rus janna 624] [LN lethanoco 81 of 1 rcent] [I	[OR.] [OR.] [SP.]	METHAN (Q58370 (CL:Ba (EC:2 39) [AC jannasc the co)] [RE: NT Length:	OCOCCUS Cillus .2.1.2 :U6753 hii] [Implete 8643] AA Length	S JANN swissp subti] [DB: 9:L771 DB:gen genom [DI:co	ASCHII] rot] lis 23K pir2] 17] pept-bct2] e.] [NT:si mplement]	ity
Description sp:[LN:TAL_METJA] [AC:Q58370] [GN:TAL [EC:2.2.1] [DE:TRANSALDOLASE-LIKE >pir:[LN:H64419] [AC:H64419] [PN:tr phosphoprotein orfU] [OR:Methanococc [MP:REV892437-891784] >gp:[GI:g1591 [PN:transaldolase] [GN:MJ0960] [OR:M [DE:Methanococcus jannaschii section SP:P19669 PID:853766 GB:AL009126 per ORF Name AT7503000993_12894378_f2_347 Description NO-HIT	AL:MJ0960 PROTEIN, ransaldol rus janna 624] [LM lethanoco 1 81 of 1 rcent] [I	[OR.] [OR.] [SP.]	METHAN :Q58370 [CL:Ba [EC:2 39] [AC jannasc the co 0] [RE: NT Length:	OCOCCUS Cillus .2.1.2 :U6753 hii] [Implete 8643] AA Length	S JANN swissp subti] [DB: 9:L771 DB:gen genom [DI:co	ASCHII] rot] lis 23K pir2] 17] pept-bct2] e.] [NT:si mplement] probabil	ity
Description sp:[LN:TAL_METJA] [AC:Q58370] [GN:TAL [EC:2.2.1] [DE:TRANSALDOLASE-LIKE >pir:[LN:H64419] [AC:H64419] [PN:tr phosphoprotein orfU] [OR:Methanococc [MP:REV892437-891784] >gp:[GI:g1591 [PN:transaldolase] [GN:MJ0960] [OR:M [DE:Methanococcus jannaschii section SP:P19669 PID:853766 GB:AL009126 per ORF Name A17503000993_12894378_f2_347 Description NO-HIT ORF Name	AL:MJ0960 PROTEIN, ransaldol rus janna .624] [LN lethanocc 1 81 of 1 rcent] [I NTID 2142 NTID	[OR] [OR] [SP] [SP] [SP] [SP] [SP] [SP] [SP] [SP	METHAN :Q58370 [CL:Ba [EC:2 39] [AC jannasc the co 0] [RE: NT Length:	OCOCCUS CILLUS C	S JANN swissp subti] [DB: 9:L771 DB:gen genom [DI:co	ASCHII] rot] lis 23K pir2] 17] pept-bct2] e.] [NT:si mplement] probabil	ity
Description sp:[LN:TAL_METJA] [AC:Q58370] [GN:TAL [EC:2.2.1] [DE:TRANSALDOLASE-LIKE >pir:[LN:H64419] [AC:H64419] [PN:tr phosphoprotein orfU] [OR:Methanococc [MP:REV892437-891784] >gp:[GI:g1591 [PN:transaldolase] [GN:MJ0960] [OR:M [DE:Methanococcus jannaschii section SP:P19669 PID:853766 GB:AL009126 per ORF Name AT7503000993_12894378_f2_347 Description NO-HIT ORF Name	AL:MJ0960 PROTEIN, ransaldol rus janna .624] [LN lethanocc 1 81 of 1 rcent] [I NTID 2142 NTID	[OR] [OR] [SP] [SP] [SP] [SP] [SP] [SP] [SP] [SP	METHAN :Q58370 [CL:Ba [EC:2 39] [AC jannasc the co 0] [RE: NT Length:	OCOCCUS CILLUS C	S JANN swissp subti] [DB: 9:L771 DB:gen genom [DI:co	ASCHII] rot] lis 23K pir2] 17] pept-bct2] e.] [NT:si mplement] probabil	ity
Description sp:[LN:TAL_METJA] [AC:Q58370] [GN:TAL [EC:2.2.1] [DE:TRANSALDOLASE-LIKE >pir:[LN:H64419] [AC:H64419] [PN:tr phosphoprotein orfU] [OR:Methanococc [MP:REV892437-891784] >gp:[GI:g1591 [PN:transaldolase] [GN:MJ0960] [OR:M [DE:Methanococcus jannaschii section SP:P19669 PID:853766 GB:AL009126 per ORF Name A17503000993_12894378_f2_347 Description NO-HIT ORF Name A17503000993_13001537_f3_444 Description	AL:MJ0960 PROTEIN, ransaldol rus janna .624] [LN lethanocc 1 81 of 1 rcent] [I NTID 2142 NTID	[OR] [OR] [SP] [ASE,] [SP] [ASE,] [SP] [ASE] [AS	METHAN (Q58370 [CL:Ba [EC:239] [AC jannasc the co] [RE: NT Length Length Length Length	OCOCCUS Collus C	S JANN swissp subti] [DB: 9:L771 DB:gen genom [DI:co	ASCHII] rot] lis 23K pir2] 17] pept-bct2] e.] [NT:si mplement] probabil	ity

ORF Name	NTID	AAID	<u>NT</u> Length	<u>AA</u> Length	core	probability
A17503000993_1352042_c2_777	2145	5917	180	59		
Description						
NO-HIT						
ORF Name	NTID	AAID	<u>NT</u> Lengthl	AA Length S	core	probability
A17503000993_13678452_c1_628	2146	5918	1263	420	438 2	.9e-41
Description						
sp:[LN:ECSB_BACSU] [AC:P55340] [GN:ESP:P55340] [DB:swissprot] >pir:[LN protein) ecsB] [GN:ecsB] [OR:Bacil:LN:BSECSABCG] [AC:X87807] [PN:hyposubtilis] [DB:genpept-bct1] [DE:B.st[LE:984] [RE:2210] [DI:direct] >gp:[AC:Z99109:AL009126] [PN:ABC transpeboth components of the protein] [OR subtilis complete genome (section 6 name: prsT, yhaC] [SP:P55340] [LE:76 >gp:[GI:e324951:g2226114] [LN:BSY140] [OR:Bacillus subtilis] [DB:genpept-lglyB-prsA region.] [NT:Identified as [LE:1538] [RE:2764] [DI:complement]	:G69619] lus subt: thetical ubtilis ([GI:e118: orter (me :Bacillus of 21): 8149] [RI 077] [AC	[AC:Gilis] ECSB ECSA, 3007:gembran s subt from E:7937 :Y1407 E:Baci	69619] [DB:pir: protein ecsB, a: 2633341 e prote ilis] [] 999501 : 5] [DI: 7] [PN:] llus sul	[PN:AB 2] >gp:] [GN:e nd ecsC] [LN:B in)] [G DB:genp to12099 direct] Hypothe bitlis	C tran [GI:e1 csB] [genes SUB000 N:ecsE ept-bc 40.] [tical 10.6 K	asporter (membrane .83450:g1177254] [OR:Bacillus .] [SP:P55340] [S] [FN:regulates [t1] [DE:Bacillus [NT:alternate gene [protein] [GN:yhaC] [tb chromosomal DNA:
ORF Name	NTID	AAID	NT LengthI	AA Length s	core	probability
AI7503000993 13722338 c3 943	2147	5919			505 2	.3e-48
Description] []		الـــال			
pir:[LN:F69992] [AC:F69992] [PN:th:peroxidase] [OR:Bacillus subtilis] [AC:Z99119:AL009126] [GN:ytgI] [FN:totalis] [DE:Bacillus subtilis complete genor [NT:similar to thiol peroxidase] [LN:peroxidase] [LN:peroxidase] [AC:gp:[GI:g2293238] [LN:AF008220] [AC:subtilis] [DB:genpept-bct2] [DE:BacilnT:similarity to tagD protein from	[DB:pir2] unknown] me (sect: E:18998] :AF008220 illus sub	>gp: OR:B ion 16 RE:1 D] PN	[GI:ell: acillus of 21) 9501] [I :YtgI] rrnB-di	85822:g subtil : from DI:comp [GN:ytg naB gen	263543 is] [D 299777 lement I] [OR omic r	3] [LN:BSUB0016] B:genpept-bct1] 1to 3213410.]] ::Bacillus egion.]
ORF Name	NTID	AAID	<u>NT</u> Lengthi	AA Length	core	probability
AI7503000993_13723318_c3_945	2148	5920	1149	382	1230 3	.4e-125
Description						
pir:[LN:JE0388] [AC:JE0388] [PN:ala [EC:1.4.1.1] [DB:pir3] >gp:[GI:d104] [PN:alanine dehydrogenase] [GN:alad]	1279:g480	3749]	[LN:ABO	013821]	[AC:A	B013821]

aerogenes DNA] [DB:genpept-bct1] [EC:1.4.1.1] [DE:Enterobacter aerogenes aladh gene for alanine dehydrogenase, complete cds.] [LE:174] [RE:1307] [DI:direct]

ORF Name	NTID	AAID	NT AA LengthLengt	h score	probability	
AI7503000993_13787912_c3_891	2149	5921	1524 507		1.7e-05	\neg
Description pir: [LN:D64924] [AC:D64924] [PN:hyp [DB:pir2] >gp:[GI:g1549279] [LN:ECU6 [DB:genpept-bct1] [DE:Escherichia co [NT:hypothetical protein] [LE:4535] [LN:AE000262] [AC:AE000262:U00096] Unknown] [OR:Escherichia coli] [DB:g section 152 of 400 of the completege identical (6 gaps)] [LE:872] [RE:247	58703] oli K-12 [RE:613 [PN:orf, genpept- enome.]	[AC:U68 2 MG165 39] [DI hypot -bct2] [NT:o5	703] [OR:Esc 5 genome, ri :direct] >gp hetical prot [DE:Escheric 34; This 534	herichi bC-pykF :[GI:g1 ein] [G	.a coli] 7 region.] .787957] &N:b1668] [FN:or .i K-12 MG1655	-
ORF Name	NTID	AAID	NT AA LengthLengt	h score	probability	
AI7503000993_13852187_f1_136	2150	5922	186 61		4.5e-06	٦
Description pir: [LN:S77632] [AC:S77632:S52761] [OR:Staphylococcus aureus phage phi- [AC:X82312] [PN:integrase] [GN:int] [OR:Bacteriophage phi-13] [DB:genper] [LE:461] [RE:1498] [DI:direct]	13] [DE [FN:int	3:pir2] :egrati	>gp:[GI:g75 on of phi-13 cteriophage	8229] [in S.a	LN:PHI13INT] ureus genome]]
ORF Name	NTID	<u>AAID</u>	NT AA Length Lengt	h score	probability	
AT7503000993_13939027_f3_560	2151	5923	186 61	Ī		
Description NO-HIT						
ORF Name	NTID	AAID	NT AA LengthLengt	h score	probability	
AT7503000993_14251933_f2_239	2152	5924	663 220		1.5e-81	\neg
Description pir: [LN:A37146] [AC:A37146:A44901:S1 S4:ribosomal protein BS4 (rpsD)] [GN [OR:Bacillus subtilis] [DB:pir2] >gp [PN:ribosomal protein S4] [GN:rpsD] isolate BR151) DNA] [DB:genpept-bct1 complete cds and tyrosyl tRNAsynthet [DI:direct] >gp:[GI:e1185839:g263545] protein S4 (BS4)] [GN:rpsD] [OR:Baci subtilis complete genome (section 16 [LE:37035] [RE:37637] [DI:direct] >g [PN:ribosomal protein S4] [GN:rpsD] [DE:Bacillus subtilis rrnB-dnaB geno [DI:complement]	I:rpsD] D:[GI:g1 [OR:Bac] [DE:E ase (ty [O] [LN:] llus su [of 21) [p:[GI:g [OR:Bac	[CL:E .43467] :illus 3.subti rrs) ge BSUB00 ubtilis : from [229331	scherichia c [LN:BACRPSD subtilis] [S lis ribosoma ne, 3' end.] 16] [AC:Z991] [DB:genpep 2997771to 3 9] [LN:AF008 subtilis] [D [LE:142790]	oli rib] [AC:M R:B.sub l prote [LE:75 19:AL00 t-bct1] 213410. 220] [A B:genpe	cosomal protein (159358:M34718) (159358:M34718	68,
ORF Name	NTID	AAID	NT AA LengthLength	<u>score</u>	probability	
AI7503000993_14460877_c3_963	2153	5925	138 45	131	2.8e-08	
Description gp:[GI:g1022726] [LN:SHU35635] [AC:U haemolyticus] [SR:Staphylococcus hae [DE:Staphylococcus haemolyticus IS12 [LE:1101] [RE:1922] [DI:complement]	molytic	us str	ain=Y176] [D	B:genpe	pt-bct1]	

		NT AA
ORF Name	NTID	AAID LengthLength score probability
AI7503000993_14460882_c1_649	2154	5926 177 58 227 6.6e-19
Description gp:[GI:g1022726] [LN:SHU35635] [AC:U haemolyticus] [SR:Staphylococcus hae [DE:Staphylococcus haemolyticus IS12 [LE:1101] [RE:1922] [DI:complement]	molytic	
ORF Name	NTID	AAID NT AA score probability
A17503000993_14460932_c1_612 Description	2155	5927 177 58 211 3.3e-17
gp:[GI:g1022726] [LN:SHU35635] [AC:Uhaemolyticus] [SR:Staphylococcus hae	molytic	
ORF Name	NTID	AAID NT AA score probability
AI7503000993_14460932_f2_226	2156	5928 168 55 196 1.3e-15
Description gp:[GI:g1022726] [LN:SHU35635] [AC:Uhaemolyticus] [SR:Staphylococcus haemolyticus IS12 [LE:1101] [RE:1922] [DI:complement]	molytic	- -
ORF Name	NTID	AAID NT AA score probability
AI7503000993_14656327_c2_751	2157	5929 624 207 799 1.6e-79
	pt-bct2	:U76550] [PN:CadD] [GN:cadD]] [DE:Staphylococcus aureus plasmid pRW001 low level cadmium resistance] [LE:2328]
ORF Name	NTID	AAID NT AA score probability
AI7503000993_14658152_£3_589	2158	5930 156 51
Description NO-HIT		
ORF Name	NTID	AAID NT AA score probability
AI7503000993_14742937_c2_774	2159	5931 231 76
Description		

ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000993_14877316_f1_64	2160	5932	407 5.5e-38
Description	L	J L	
membrane protein b1832] [CL:hypother [DB:pir2] >gp:[GI:d1016364:g173647] YKL340] [OR:Escherichia coli] [SR:Escherichia coli] [SR:Escherichia coli] [SR:Escherichia coli] [SR:Escherichia coli] [SR:Escherichia coli] [SR:Escherichia coli] [M:CORF_ID:0335#13; similated [DI:complement] >gp:[GI:d1016369:g1] [GN:YKL069W, YKL340] [OR:Escherichia colone_lib:Kohara lambda minise] [DI:#336(41.2-41.6 min.).] [NT:ORF_ID:08132] [RE:3299] [DI:complement] >gp:[GI:ghypothetical protein] [GN:b1832] [DB:genpept-bct2] [DE:Escherichia colone]	N PRC-PP sprot] > etical p 3] [LN:D Escheric DE:E.col r to [Sw 1736479] ia coli] B:genpep 0335#13; g1788136 FN:orf; coli K-1	HA INTERPRETATION TO THE PROPERTY OF THE PROPE	ERGENIC REGION] N:H64944] [AC:H64944] [PN:probable YKL069w] [OR:Escherichia coli] [AC:D90826:AB001340] [GN:YKL069W, Li (strain:K12) DNA, clone_lib:Kohara mic DNA, Kohara clone #335(40.9-41.3 L Accession] [LE:16795] [RE:17346] B0827] [AC:D90827:AB001340] Scherichia coli (strain:K12) DNA, [DE:E.coli genomic DNA, Kohara clone ar to [SwissProt Accession] [LE:2748] AE000277] [AC:AE000277:U00096] [PN:orf, Li [OR:Escherichia coli]
ORF Name	NTID	AAID	NT AA LengthLength
A17503000993_14882928_c3_923	2161	5933	1098 365 1279 2.2e-130
Description		J	
gp:[GI:e220317:g1177684] [LN:SXCCP] [OR:Staphylococcus xylosus] [DB:genacuC and acuA genes.] [LE:<1] [RE:	npept-bc	t1] [E0	C:5.4.99.5] [DE:S.xylosus aroA, ccpA,
ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000993_15650303_c3_882	2162	5934	207 68
Description		<i>-</i>	
NO-HIT			
ORF Name	NTID	AAID	NT AA LengthLength score probability
A17503000993_163151_c3_886	2163	5935	1248 415 1800 1.3e-185
Description			
sp:[LN:METK_STAAU] [AC:P50307] [GN [DE:ADENOSYLTRANSFERASE) (ADOMET SY >gp:[GI:g1020317] [LN:SAU36379] [AC [FN:catalyses the synthesis of SAM] [DE:Staphylococcus aureus S-adenosy synthetase] [LE:212] [RE:1405] [DI	YNTHETASI C:U36379] [OR:Staylmethion	E)] [SI] [PN:S aphyloc	P:P50307] [DB:swissprot] G-adenosylmethionine synthetase]
ORF Name	NTID	AAID	NT AA score probability
AI7503000993_16533442_c2_738	2164	5936	522 173 546 1.0e-52
Description		J	
	DB:ger	npept-k	pR] [GN:cspR] [OR:Bacillus subtilis] oct1] [DE:Bacillus subtilis methylase homolog] [LE:573] [RE:1046]

A17503000993_16586012_f2_308 2165 5937 588 195 456 3.5e-43
Description
pir:[LN:E69999] [AC:E69999] [PN:hypothetical protein ytqB] [GN:ytqB] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:e1185922:g2635533] [LN:BSUB0016] [AC:Z99119:AL009126] [GN:ytqB] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 16 of 21): from 2997771to 3213410.] [LE:122275] [RE:122859] [DI:direct] >gp:[GI:g2293301] [LN:AF008220] [AC:AF008220] [PN:YtqB] [GN:ytqB] [OR:Bacillus subtilis] [DB:genpept-bct2] [DE:Bacillus subtilis rrnB-dnaB genomic region.] [LE:57568] [RE:58152] [DI:complement]
ORF Name NTID AAID NT AA score probability
AI7503000993_165908_f2_353
Description sp:[LN:YHGC_BACSU] [AC:P38049] [GN:YHGC] [OR:BACILLUS SUBTILIS] [DE:HYPOTHETICAL 18.8 KD PROTEIN IN ECSC-PBPF INTERGENIC REGION] [SP:P38049] [DB:swissprot] >pir:[LN:B40614] [AC:B40614:F69832] [PN:conserved hypothetical protein yhgC:hypothetical protein X (pbpF 5' region)] [GN:yhgC] [OR:Bacillus subtilis] [DB:pir2] >gp:[GI:g304160] [LN:BACPBPF] [AC:L10630] [OR:Bacillus subtilis] [SR:Bacillus subtilis (strain W168) DNA] [DB:genpept-bct1] [DE:Bacillus subtilis penicillin-binding protein (pbpF) gene, 5' end.] [NT:product unknown] [LE:247] [RE:747] [DI:complement] >gp:[GI:e1183012:g2633346] [LN:BSUB0006] [AC:Z99109:AL009126] [GN:yhgC] [FN:unknown] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis complete genome (section 6 of 21): from 999501 to1209940.] [NT:alternate gene name: yixC; similar to hypothetical] [SP:P38049] [LE:83202] [RE:83702] [DI:complement] >gp:[GI:e325006:g2226228] [LN:BSY14083] [AC:Y14083] [PN:Hypothetical protein] [GN:yixC] [OR:Bacillus subtilis] [DB:genpept-bct1] [DE:Bacillus subtilis chromosomal DNA, region 76-78 degrees: betweenglyB-aprE.] [NT:See Swiss Prot P38049; YIXC_BACSU] [SP:P38049] [LE:3061] [RE:3561] [DI:complement]
NTID AAID NT AA score probability A17503000993_16835053_c2_758 2167 5939 168 55
Description NO-HIT
ORF Name NTID AAID LengthLength score probability A17503000993_16994043_c3_933
Description pir:[LN:E69106] [AC:E69106] [PN:hypothetical protein MTH1793] [GN:MTH1793] [OR:Methanobacterium thermoautotrophicum] [DB:pir2] >gp:[GI:g2622925] [LN:AE000934] [AC:AE000934:AE000666] [PN:unknown] [GN:MTH1793] [OR:Methanobacterium thermoautotrophicum] [DB:genpept-bct2] [DE:Methanobacterium thermoautotrophicum from bases 1640298 to 1655421(section 140 of 148) of the complete genome.] [NT:Function Code:14.00 - Unknown, ; similar to,] [LE:89] [RE:790] [DI:complement]
<u>NTID AAID NT AA</u> <u>LengthLength</u> score probability A17503000993_189437_f1_34 2169
Description NO-HIT

NTID

ORF Name

AAID NT AA score probability